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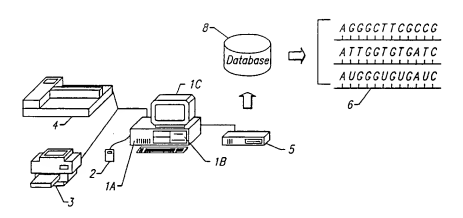
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(54) Title: OLIGOPROBE DESIGNSTATIONS: A COMPUTERIZED METHOD FOR DESIGNING OPTIMAL OLIGON-UCLEOTIDE PROBES AND PRIMERS



(57) Abstract

There is disclosed herein an invention which relates to the fields of genetic engineering, microbiology, and computer science, that allows a user, whether a molecular biologist or a clinical diagnostician, to calculate and design extremely specific oligonucleotide sequences for DNA and mRNA hybridization procedures. The sequences designed with this invention may be used for medical diagnostic kits, DNA indentification, and potentially continuous monitoring of metabolic processes in human beings. The key features design oligonucleotide sequences based on the GenBank database of DNA and mRNA sequences and examine candidate sequences for specificity or commonality with respect to a user-selected experimental preparation. Two models are available: a Mismatch Model, that employs hashing and continuous seed filtration, and an H-site Model, that analyzes candidate sequences for their binding specificity relative to some known set of mRNA or DNA sequences. The preferred embodiment of this computerized design tool is written in the Borland R C++ language and runs under Microsoft R Windows TM on IBM R compatible personal computers.

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OLIGOPROBE DESIGNSTATION: A COMPUTERIZED METHOD FOR DESIGNING OPTIMAL OLIGONUCLEOTIDE PROBES AND PRIMERS

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BACKGROUND OF THE INVENTION

This invention relates to the fields of genetic engineering, microbiology, and computer science, and more specifically to an invention that helps the user, whether they be a molecular biologist or a clinical diagnostician, to calculate and design extremely accurate oligonucleotide sequences for use as probes, for example for DNA and mRNA hybridization procedures, or as primers, for example for DNA amplification and extension using the polymerase chain reaction (PCR). In the following description, the design of probes has been discussed.

The oligonucleotide probes designed with this invention may be used to test for the presence of precursors of specific proteins in living tissues, or may be used for medical diagnostic kits, DNA identification, and potentially continuous monitoring of metabolic processes in human beings. The present implementation of this computerized design tool runs under Microsoft ® Windows W. 3.1 (made by Microsoft Corporation of Redmond, Washington) on IBM ® compatible personal computers (PC's).

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Although any methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, the preferred methods and materials are now described. All publications mentioned hereunder are incorporated herein by reference.

To isolate a specific gene for any particular purpose, a researcher first has to have some idea of what he or she is looking for. To do this, the researcher needs to have a probe, which acts like a molecular hook that can identify and latch onto (i.e., bind to or hybridize with) the desired gene in a crowd of many other genes. A researcher who can obtain an entire strand of mRNA can eventually find the gene from which it was copied, using complementary DNA (cDNA, which is a cloned equivalent

to RNA and somewhat equivalent to mRNA) as a probe to search through the great mass of genetic material and locate the desired original gene. cDNA essentially is manufactured or non-naturally occurring DNA from which all of the nonessential DNA has been removed. cDNA allows the researcher to concentrate entirely on the important portions of the gene being examined. The nonessential DNA regions are easy to recognize because when the gene is translated into protein, these regions do not wind up reflected in the protein sequence. These regions are called introns, or intervening regions. mRNA has no introns because they have been "spliced" out of the mRNA before translation. Thus, mRNA and cDNA contain only the essential information from a gene (called the exons). cDNA is the equivalent of mRNA with a complementary sequence, only the exons are present. cDNA may be produced by reverse transcription of mRNA.

The procedure of using cDNA from known mRNA as a probe to search through genetic material and locate the original gene is called molecular hybridization, and is currently one method of identifying specific genes. However, this method is less than perfect, can be extremely time consuming, and often is not even feasible because the researcher actually has to have an entire strand of cDNA from the desired gene before he or she can attempt to use this cDNA to locate and identify the particular gene. Thus, it is something of a circular problem. If the researcher cannot obtain an entire strand of mRNA or cDNA from the desired gene, then he or she must somehow design a probe from scratch to be used to identify that gene.

Oligonucleotide probes (that is, probes made up of a small number of nucleotides, such as 17 to 100), are increasingly being used to identify specific genes from genomic or cDNA libraries when the partial amino acid sequences is known. (von Heijne 1987, Ref. 15). This is a second method of determining a proper probe. Although the present implementation of this invention does not deal with cases in which the proteins have been sequenced, but rather only the DNA or mRNA, it is possible that this invention or a future implementation of it might be used with protein sequences. Such probes can also be used as primers which, when annealed to mRNAs, can be selectively extended into cDNAs. (von Heijne 1987, Ref. 15).

Because of these situations, the problem that the researcher faces is to discover or design a probe or mixture of probes that maximizes the researchers chances of successful hybridization while at the same time minimizing the amount of time and money that has to be spent on discovering or designing the probes. (von Heijne 1987,

Ref. 15). Researchers in the field have determined that computer analysis can greatly expedite and simplify the search for optimal probe sequences. (von Heijne 1987, Ref. 15). However, all of the search strategies known to the present inventors are time consuming (both CPU and user time) and may be somewhat inaccurate. As stated in von Heijne, "a true optimization of the probe in terms not only of degeneracy but in terms of length, codon usage, Guanine-Cytosine (GC) avoidance, and expected signal-to-noise ratio (hybridization to target over background) is a fairly complex problem, however, and does not seem to have been automated so far." (von Heijne 1987, Ref. 15). Various search strategies known and used in the field to identify and design probes are outlined in the following sources: Lewis (1986, Ref. 9), Raupach (1984, Ref. 11), Yang et al. (1984, Ref. 16), and Martin and Castro (1984, Ref. 10).

In the simplest version of a protein-related search strategy, the search procedure is limited to finding a set of probes of given lengths with the least possible degeneracy simply by scanning the amino acid sequence and noting the number of alternative codons in the corresponding oligonucleotide as the scan moves along the chain of nucleotides. (Lewis 1986). The researcher can also include codon usage statistics (because more than one codon can translate to the same amino acid), which would attach a probability-of-occurrence value to each probe. (Raupach 1984, Ref. 11).

A more advanced algorithm would allow the researcher to specify the way in which he or she plans to synthesize the probes (for example, by adding monomers or mixtures of monomers). It would also be easy for a researcher to add a rough estimate of the disassociation (or melting) temperatures of each probe to a program such as this.

One way to solve the problem of finding local similarities between two proteins being compared that has been discussed in the relevant literature is to use list-sorting or hashing routines. (von Heijne 1987, Ref. 15). These routines are based on the construction of a list or lookup table of k-letter words or k-tuples (i.e., all possible dior trinucleotides), and the positions where they appear in the sequences being compared. This method is employed in some of the most extensively used "fast search" programs (see examples identified in von Heijne 1987, Ref. 15).

Two general methods of designing probes are common in the field, depending upon whether the researcher is trying to design a common probe or a specific probe. Common probes attempt to find common or consensus sequences among various species and among family genes. The first step in designing such a probe is to find the genes of interest. This may be done by performing a keyword or homology search against the

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GenBank (a genome database available from IntelliGenics of Mountain View, CA) or a keyword search against MEDLINE (the database currently available from the U.S. National Library of Medicine under the data access system known as Dialog of Dialog Information Service, Inc., Palo Alto, CA) or by performing a homology analysis between one of the genes of interest and whole GenBank sequences. The next step is to retrieve all of the relevant genes of interest. In the third step, multiple alignment analysis can be done using a commercially available software package such as DNASIS (from Hitachi Software of Brisbane, California), which is an autoconnect program. In this step, the computer identifies which nucleotides are common among the requested sequences:

* = common among A1, A2, and A3

Alternatively, after homology analyses between two sequences are carried out, data from the multiple homology analyses can be combined. The researcher then manually has to find the common or consensus region:

* = common among A1, A2, and A3

Next, the researcher would input the sequence of the common region into the program and then analyze the secondary structure (i.e., the stacking site and the hairpin structure). After this, the researcher manually would select several candidate probes (from five to ten) which contain the minimal hairpin structure and specific length according to the user's interest. A hairpin is an area in which a probe has "folded back" and one portion of the probe has hybridized with another portion of the same probe. The researcher would then perform a homology analysis between each candidate probe and all sequences in the GenBank to find all possible cross-hybridizable genes. Lastly,

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the researcher manually would decide which is the best candidate probe by determining which probe is highly homologous among the group of interest, but quite different from other unrelated sequences in the GenBank.

The conventional methods for designing common oligonucleotide probes using currently available computer software have at least five problems: (1) they involve time consuming multiple processes; (2) it is difficult to control a significant variable, the melting temperature Tm of the oligonucleotide probes; (3) the methods do not recognize exons and introns and differentiate (thereby making it possible to have a designed probe that is identical to unrelated mRNA sequences); (4) the methods may miss short pieces of identical sequences; and (5) it is difficult to recognize multiple pieces of identical sequences in the gene.

The second method of designing probes that is common in the field involves designing specific probes. Specific probes attempt to find unique sequences among various species and among family genes and among published sequences in the GenBank. A specific probe is a probe that hybridizes with only one particular gene, thereby identifying the presence of that gene for the researcher. The procedure involves first finding the genes of interest (by performing a keyword search against the GenBank or against MEDLINE) and then retrieving all of the relevant genes of interest. A manual homology analysis between the gene of interest and whole sequences in the GenBank can be performed to find common and unique regions.



Next, the researcher would input the sequence of the unique region into the program and then analyze the secondary structure. After this, the researcher would manually select several candidate probes which contain the minimal hairpin structure and specific length according to the user's interest. The researcher would then perform a homology analysis between each candidate probe and all sequences in the GenBank to find all possible cross-hybridizable genes. Lastly, the researcher manually would decide which is the best candidate probe by determining which probe does not have identical sequences in unrelated sequences in the GenBank.

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All of the conventional methods for designing specific oligonucleotide probes known to the inventors using currently available computer software have at least four problems: (1) they involve time consuming multiple processes; (2) it is difficult to control the melting temperature Tm of the oligonucleotide probes; (3) the methods do not allow for quantification of uniqueness; and (4) there is no guarantee that the method will design the best possible probe.

None of the methods discussed in the literature discloses a system that may be used to design both common probes <u>and</u> extremely specific probes, especially a method that minimizes user and CPU time and is exceptionally accurate.

Programs currently used for rapid database similarity searches use either hashing strategies or statistical strategies. The hashing strategy is now being used for the detection of relatively short regions of similarity, while the statistical strategy is now being used for the detection of weaker and longer similarity regions. The Mismatch Model of this invention can be used for very strong similarity searches with running times faster than current hashing strategies.

The basic technologies behind the Mismatch Model used in this invention are hashing and continuous seed filtration, each general technology being known in the public domain and having been previously applied separately to non-genetic applications. To the best of the inventors' knowledge, these methods, used together, have never been suggested in other studies on optimal probe selection. The inventors' methods have a program performance of tens of seconds (CPU + I/O time) with a 1000 nucleotide query and all mammalian DNA on a SPARC station, and are even faster on the more common personal computer proposed herein.

The H-Site Model of this invention likewise is unique in that it offers a multitude of information on selected probes and original and distinctive means of visualizing, analyzing and selecting among candidate probes designed with the invention. Candidate probes are analyzed using the H-Site Model for their binding specificity relative to some known set of mRNA or DNA sequences, collected in a database such as the GenBank database. The first step involves selection of candidate probes at some or all the positions along a given target. Next, a melting temperature model is selected, and an accounting is made of how many false hybridizations each candidate probe will produce and what the melting temperature of each will be. Lastly, the results are presented to the researcher along with a unique set of tools for visualizing, analyzing and selecting among the candidate probes.

This invention is both much faster and much more accurate than the methods that are currently in use. It is unique because it is the only method that can find not only the most specific and unique sequence, but also the common sequences. Further, it allows the user to perform many types of analysis on the candidate probes, in addition to comparing those probes in various ways to the target sequences and to each other.

Therefore, it is the object of this invention to provide a practical and user-friendly system that will allow a researcher to design both specific and common oligonucleotide probes, and to do this in less time and with much more accuracy than currently done. For example, the current version of the GenBank contains over ninety (90) million nucleotides. It is thought that the human genome alone consists of three billion base pairs, and scientists have so far managed to decode the base sequence of only about 500 human genes, less than one percent of the total. Currently available searching strategies are limited in how many of the GenBank's sequences can be accessed and successfully searched, and how convenient and feasible such a search would be (in terms of both computer processor and human user time). It is also an object of this invention to allow the user to be able to run the program on more readily available and far less expensive computer hardware (i.e., a PC rather than a mainframe). This invention will remove those limits and allow genetic research to take a giant leap forward.

These and other advantages and objects of this invention will become apparent from the following detailed descriptions, drawings, and appended claims.

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BRIEF DESCRIPTION OF THE INVENTION

There is disclosed herein a system which allows the user to calculate and design extremely accurate oligonucleotide probes for DNA and mRNA hybridization procedures. The invention runs under Microsoft ® Windows on IBM ® compatible personal computers (PC's). Its key features design oligonucleotide probes based on the GenBank database of DNA and mRNA sequences and examine probes for specificity or commonality with respect to a user-selected experimental preparation of gene sequences. Hybridization strength between a probe and a subsequence of DNA or mRNA can be estimated through a hybridization strength model. Quantitatively, hybridization strength is given as the melting temperature Tm. Currently, two hybridization strength models are supported by this invention: 1) the Mismatch Model and 2) the H-Site Model. The user is allowed to select from the following calculations for each probe, results of which are available for display and analysis: 1) Sequence, Melting Temperature (Tm) and Hairpin characteristics; 2) Hybridization with other species within the preparation mixture; and (3) Location and Tm for the strongest hybridizations. The results of the invention's calculations are then displayed on the Mitsuhashi Probe Selection Diagram (MPSD), which is a graphic display of all of the hybridizations of probes for the target mRNA with all sequences in the preparation.

The Main Dialog Window of the present implementation of this invention controls all user-definable settings. The user is offered a number of options at this window. The File option allows the user to print, print in color, save selected probes, and exit the program. The Preparation option allows the user to open and create preparation (PRP) files. The Models option allows the user to chose between the two hybridization models currently supported by the invention: 1) the H-Site Model and 2) the Mismatch Model. If the user selects the H-Site Model option, the user normally sets the following model parameters: 1) the melting temperature Tm for which probes are being designed (i.e., the melting temperature that corresponds to a particular experiment or condition the user desires to simulate); and 2) the nucleation threshold, which is the number of base pairs constituting a nucleation site. If the user selects the Mismatch Model option, the user normally sets the following model parameters: 1) probe length, which is the number of bases in probes to be considered; and 2) mismatch N, which is the maximum number of mismatches constituting a hybridization.

The Mismatch Model program is used to design DNA and mRNA probes, utilizing sequence database information from sources such as GenBank and other

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databases with similar file formats. In the Mismatch Model, hybridization strength is related only to the number of base pair mismatches between a probe and its binding site. Generally, the more mismatches a user allows, the more probes will be found. The Mismatch Model does not take into account the Guanine-Cytosine (GC) content of candidate probes, as does the H-Site Model, discussed below, so there is no reflection or indication of the probe's binding strength. The basic technologies employed by this model are hashing and continuous seed filtration. Hashing involves the application of an algorithm or process to the records in a set of data to obtain a symmetric grouping of the records. When using an indexed set of data, hashing is the process of transforming a record key to an index value for storing and retrieving a record. Rosenberg (1984, Ref. 12)). The concept of continuous seed filtration is discussed in detail below.

The essence of the Mismatch Model is a fast process for doing exact and inexact matching between DNA and mRNA sequences to support the Mitsuhashi Probe Selection Diagram (MPSD) and other types of analysis discussed above. The process used by the Mismatch Model is the Waterman-Pevzner Algorithm (the WPALG, which is named for two of the inventors), which is a computer-based probe selection process. Essentially, this is a combination of new and improved pattern matching processes. See Hume and Sunday (1991, Ref. 4), Landau et al (1986-1990, Refs. 6, 7, 8), Grossi and Luccio (1989, Ref. 3), and Ukkonen (1982, Ref. 14).

There are three principal programs that make up the Mismatch Model in this implementation of the invention. The first is designated by the inventors as "k_diff." WPALG is used in k_diff to find all locations of matches of length greater than or equal to one (1) (length is user-specified) with less than or equal to k number of mismatches (k is also user-specified) between the two sequences. If a candidate oligonucleotide probe fails to match that well, it is considered unique. k_diff uses hashing and continuous seed filtration, and looks for homologs in GenBank and other databases with similar file formats. The technique of continuous seed filtration allows for much more efficient searching than previously implemented techniques. A seed is defined in this invention to be a subsequence of length equal to the longest exact match in the worst case scenario. For example, suppose the user selects a probe length (1) of 18, with 2 or fewer mismatches (k). If a match exists with 2 mismatches, then there must be a perfectly matching subsequence of length equal to 6. Once the seed length has been determined, the Mismatch Model looks at all substrings of that seed length (in this

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example, that seed length would be 6), finds the perfectly matched base pair subsequence of length equals 6, and then looks to see if this subsequence extends to a sequence of length equal to the user selected probe length (i.e., 20 in this example). If so, a candidate probe has been found that meets the user's criteria.

Where the seed size is large, the program allocates a relatively large amount of memory for the hash table. This invention has an option that allows memory allocation for GenBank entries just once at the beginning of the program, instead of reallocating memory for each GenBank entry. This reduces input time for GenBank entries by as much as a factor of two (2), but the user needs to know the maximum GenBank entry size in advance to do this.

A probe is defined to hybridize if it has k or fewer mismatches in comparison with a target sequence from the database or file searched. Otherwise, it is non-hybridizing. The hit extension time for all appropriate parameters of the Mismatch Model has been found by experimentation to be less than thirty-five (35) seconds, except in one case where the minimum probe length (1) was set to 24 and the maximum number of mismatches (k) was set to four (4), which is a situation that is never used in real gene localization experiments because the hybridization conditions are too weak.

In this invention, the second hybridization strength model is termed the H-Site Model. One aspect of the H-Site Model uses a generalization of an experimental formula in general usage. The basic formula on which this aspect of the model is built is as follows:

$$Tm = 81.5 - 16.6(log[Na]) - .63 \%(formamide) + .41 (\%(G + C)) - 600$$
 / N

In this formula, log[Na] is the sodium concentration, %(G + C) is the fraction of matched base pairs which are G-C complementary, and N is the probe length. In other words, this formula is an expression of the fact that melting temperature Tm is a function of both probe length and percent of Guanine-Cytosine (GC) content. This basic formula has been modified in this invention to account for the presence of mismatches. Each percent of mismatch reduces the melting temperature Tm by an average of 1.25 degrees (2 degrees C for an Adenine-Thymine mismatch, and 4 degrees C for a Guanine-Cytosine mismatch). This formula is, however, an approximation. The actual melting temperature might differ significantly from this approximation, especially for short probes or for probes with a relatively large number of mismatches.

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Hybridization strength in the H-Site Model is related to each of the following factors: 1) "binding region"; 2) type of mismatch (GC or AT substitution); 3) length of the probe; 4) GC content of the binding region (since GC pairs have a stronger bond than AT pairs, thus requiring a higher melting temperature); and 5) existence of a "nucleation site" (an exactly matching subsequence). The type of mismatch and the GC content of the binding region each contribute to a candidate probe's binding strength, which can be compared to other candidate probes' binding strengths to enable the user to select the optimal probe.

The fundamental assumption of the H-Site Model is that binding strength is determined by a paired subsequence of the probe-species combination, called the binding region. If the binding region contains more GC pairs than AT pairs, the binding strength will be higher since the G and C bases (connected with three bonds) form a tighter bond than the A and T bases (connected with two bonds). Thus, G and C bases, and probes that are GC rich, require a higher melting temperature Tm and subsequently form a stronger bond. In the H-Site Model, and one of its unique features, the program designs optimal probes, ideally ones that do not have any mismatches, but if there are mismatches the H-Site Model takes these into account. With this model, a candidate probe can afford to have more mismatches involving the AT bases if there are more GC bases than AT bases in the probe. This is because this model looks primarily at regions of the candidate probe and target sequence that match and does not "penalize" the probe for areas that do not match. If the mismatches are located at either or both of the ends of the binding region, this has little effect. It is much more deleterious to have mismatches in the middle of the binding region, as this will significantly lower the binding strength of the probe.

The formula cited above for Tm applies within the binding region. The length of the probe is used to calculate percentages, but all other parameters of the formula are applied to the binding region only. The H-Site Model further assumes the existence of a nucleation site, which is a region of exact match. The length of this nucleation site may be set by the user. Typically, a value of 8 to 10 base pairs is used. To complete the H-Site Model, the binding region is chosen so as to maximize the melting temperature Tm among all regions containing a nucleation site, assuming one exists (otherwise, Tm=0).

The H-Site Model is more complex than the Mismatch Model discussed above in that hybridization strength is modeled as a sum of signed contributions, with matches

generally providing positive binding energy and mismatches generally providing negative binding energy. The exact coefficients to be used depend only on the matched or mismatched pair. These coefficients may be specified by the user, although in the current version of this invention these coefficients are not explicitly user-selectable, but rather are selected to best fit the hybridization strength formulas developed by Itakura et al (1984, Ref. 5), Bolton and McCarthy (1962, Ref. 2), Benner et al (1973, Ref. 1), and Southern (1975, Ref. 13).

A unique aspect of the H-Site Model is that hybridization strength is defined to be determined by whatever the optimal binding region between the candidate probe and binding locus. This binding region is called the hybridization site, or h-site, and is selected so as to maximize overall hybridization strength, so that mismatches outside the binding region do not detract from the estimated hybridization strength. Several other unique features of the H-Site Model include the fact that it is more oriented toward RNA and especially cDNA sequences than DNA sequences, and the fact that the user has control over preparation and environmental variables. The first feature allows the user to concentrate on "meaningful" sequences, rather than having to sort through all of a DNA sequence (including the introns). The second feature allows the user to more accurately simulate laboratory conditions and more closely correspond with any experiments he or she is conducting. Further, this implementation of the invention does some preliminary preprocessing of the GenBank database to sort out and select the cDNA sequences. This is done by locating a keyword (in this case CDS) in each GenBank record, thereby eliminating any sequences containing introns.

The Mitsuhashi Probe Selection Diagram (MPSD), FIG. 4, is the third key feature of this invention, as it is a unique way of visualizing the results of the probe designing performed by the Mismatch and H-Site Models. It is a graphic display of all of the hybridizations of candidate oligonucleotide probes for the target mRNA with all sequences in the preparation. Given a gene sequence database and a target mRNA sequence, the MPSD graphically displays all of the candidate probes and their hybridization strengths with all sequences from the database. In the present implementation, each melting temperature Tm is displayed as a different color, from red (highest Tm) to blue (lowest Tm). The MPSD allows the user to see visually the number of false hybridizations at various temperatures for all candidate probes, and the sources of these false hybridizations (with a loci and sequence comparison). A locus

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may be a specific site or place, or, in the genetic sense, a locus is any of the homologous parts of a pair of chromosomes that may be occupied by allelic genes.

BRIEF DESCRIPTION OF THE DRAWING

This invention may be more clearly understood from the following detailed description and by reference to the drawing in which:

- FIG. 1 is a simplified block diagram of a computer system illustrating the overall design of this invention;
- FIG. 2 is a display screen representation of the main dialog window of this invention;
- FIG. 3 is a flow chart of the overall invention illustrating the program, and the invention's sequence and structure;
- FIG. 4 is a display screen representation of the Mitsuhashi probe selection diagram;
 - FIG. 5 is a display screen representation of the probeinfo and matchinfo window;
 - FIG. 6 is a display screen representation of the probesedit window;
 - FIG. 6a is a printout of the probesedit output file;
- FIG. 7 is a flow chart of the overall k_diff program of the Mismatch Model of this invention, including its sequence and structure;
 - FIG. 8 is a flow chart of the k diff module of this invention;
 - FIG. 9 is a flow chart of the hashing module of this invention;
 - FIG. 10 is a flow chart of the tran module of this invention;
 - FIG. 11 is a flow chart of the let dig module of this invention;
 - FIG. 12 is a flow chart of the update module of this invention;
 - FIG. 13 is a flow chart of the assembly module of this invention;
 - FIG. 14 is a flow chart of the segload module of this invention;
 - FIG. 15 is a flow chart of the read1 module of this invention;
 - FIG. 16 is a flow chart of the dig let module of this invention;
 - FIG. 17 is a flow chart of the q colour module of this invention;
 - FIG. 18 is a flow chart of the hit ext module of this invention;
 - FIG. 19 is a flow chart of the colour module of this invention;
- FIG. 20 is a printout of a sample file containing the output of the Mismatch Model program of this invention;
- FIG. 21 is a flow chart of the H-Site Model, stage I, covering the creation of a preprocessed preparation file of this invention;
- FIG. 22 is a flow chart of the H-Site Model, stage II, covering the preparation of the target sequence(s);

- FIG. 23 is a flow chart of the H-Site Model, stage III, covering the calculation of MPSD data;
- FIG. 24a is a printout of a sample file containing output of the Mismatch Model program;
- FIG. 24b is a printout of a sample file containing output of the H-Site Model program;
- FIG. 25 is a flow chart of the processing used to create the Mitsuhashi probe selection diagram (MPSD);
 - FIG. 26 is a flow chart of processing used to create the matchinfo window;
 - FIG. 27 is a printout of a sample target species file;
 - FIG. 28 is a printout of a sample preparation file.

DETAILED DESCRIPTION OF THE INVENTION

This invention is employed in the form best seen in FIG. 1. There, the combination of this invention consists of an IBM® compatible personal computer (PC), running software specific to this invention, and having access to a distributed database with the file formats found in the GenBank database and other related databases.

The preferred computer hardware capable of operating this invention involves of a system with at least the following specifications (FIG. 1): 1) an IBM © compatible PC, generally designated 1A, 1B, and 1C, with an 80486 coprocessor, running at 33 Mhz or faster; 2) 8 or more MB of RAM, 1A; 3) a hard disk 1B with at least 200 MB of storage space, but preferably 1 GB; 4) a VGA color monitor 1C with graphics capabilities of a size sufficient to display the invention's output in readable format, preferably with a resolution of 1024 x 768; and 5) a 580 MB CD ROM drive 5 (1B of FIG. 1 generally refers to the internal storage systems included in this PC, clockwise from upper right, two floppy drives, and a hard disk). Because the software of this invention preferably has a Microsoft ® Windows ™ interface, the user will also need a mouse 2, or some other type of pointing device.

The preferred embodiment of this invention would also include a laser printer 3 and/or a color plotter 4. The invention may also require a modem (which can be internal or external) if the user does not have access to the CD ROM versions of the GenBank database 8 (containing a variable number of gene sequences 6). If a modem is used, information and instructions are transmitted via telephone lines to and from the GenBank database 8. If a CD ROM drive 5 is used, the GenBank database (or specific portions of it) is stored on a number of CDs.

The computer system should have at least the Microsoft $^{\circ}$ DOS v. 5.0 operating system running Microsoft $^{\circ}$ Windows $^{\infty}$ v. 3.1. All of the programs in the preferred embodiment of the invention are written in the Borland $^{\circ}$ C++ (made by Borland International, Inc., of Scotts Valley, CA) computer language. It must be recognized that subsequently developed computers, storage systems, and languages may be adapted to utilize this invention and vice versa.

This invention is designed to enable the user to access DNA, mRNA and cDNA sequences stored either in the GenBank or in databases with similar file formats. GenBank is a distributed flat file database made up of records, each record containing a variable number of fields in ASCII file format. The stored database itself is distributed, and there is no one database management system (DBMS) common to even a majority of its users. One general format, called the line type format, is used both for

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the distributed database and for all of GenBank's internal record keeping. All data and system files and indexes for GenBank are kept in text files in this line type format.

The primary GenBank database is currently distributed in a multitude of files or divisions, each of which represents the genome of a particular species (or at least as much of it as is currently known and sequenced and publicly available). The GenBank provides a collection of nucleotide sequences as well as relevant bibliographic and biological annotation. Release 72.0 (6/92) of the GenBank CD distribution contains over 71,000 loci with a total of over ninety-two (92) million nucleotides. GenBank is distributed by IntelliGenetics, of Mountain View, CA, in cooperation with the National Center for Biotechnology Information, National Library of Medecinge, in Bethesda, MD.

1. Overall Description of the Invention

a. General Theory

The intent of this invention is to provide one or more fast processes for performing exact and inexact matching between DNA sequences to support the Mitsuhashi Probe Selection Diagram (MPSD), discussed below, and other analysis with interactive graphical analysis tools. Hybridization strength between a candidate oligonucleotide probe and a subsequence of DNA, mRNA or cDNA can be estimated through a hybridization strength model. Quantitatively, hybridization strength is given as the melting temperature Tm. Currently, two hybridization strength models are supported by the invention: 1) the Mismatch Model and 2) the H-Site Model.

b. Inputs

i. Main Dialog Window

The Main Dialog Window, FIG. 2, controls all user-definable settings. This window has a menu bar offering five options: 1) File 10; 2) Preparation 20; 3) Models 30; 4) Experiment 40; and 5) Help 50. The File 10 option allows the user to print, print in color, save selected probes, and exit the program. The Preparation 30 option allows the user to open and create preparation (PRP) files.

The Models 20 option allows the user to chose between the two hybridization models currently supported by the invention: 1) the H-Site Model 21 and 2) the Mismatch Model 25. If the user selects the H-Site Model 21 option, the left hand menu of FIG. 2C is displayed and the user sets the following model parameters: 1) the melting temperature Tm 22 for which probes are being designed (i.e., the melting temperature that corresponds to a particular experiment or condition the user desires

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to simulate); and 2) the nucleation threshold 23, which is the number of base pairs constituting a nucleation site. If the user selects the Mismatch Model 25 option, the right hand menu of FIG. 2C is displayed and the user sets the following model parameters: 1) probe length 26, which is the number of base pairs in probes to be considered; and 2) mismatch N 27, which is the maximum number of mismatches constituting a hybridization. Computation of the user's request will take longer with the H-Site Model if the threshold 23 setting is decreased and with the Mismatch Model if the number of mismatches K 27 is increased.

In addition, for both Model options the user chooses the target species 11 DNA or mRNA for which probes are being designed and the preparation 12, a file of all sequences with which hybridizations are to be calculated. A sample of a target species file is shown in FIG. 27 (humbjunx.cds), while a sample of a preparation file is shown in FIG. 28 (junmix.seq). Each of these inputs is represented by a file name and extension in general DOS format. In the target species and preparation fields, the file format follows the GenBank format, and each of the fields includes a default file extension. Pressing the "OK" button 41 of FIG. 2C will cause the processing to begin, and pressing the "Cancel" button 43 will cause it to stop.

The Experiment 40 option and the Help 50 option are expansion options not yet available in the current implementation of the invention.

c. Processing

FIG. 3 is a flow chart of the overall program, illustrating its sequence and structure. Generally, the main or "control" program of the invention basically performs overall maintenance and control functions. This program, as illustrated in FIG. 3, accomplishes the general housekeeping functions 51, such as defining global variables. The user-friendly interface 53, carries out the user-input procedures 55, the file 57 or database 59 access procedures, calling of the model program 62 or 63 selected by the user, and the user-selected report 65 or display 67, 69, 71 and 73 features. Each of these features is discussed in more detail in later sections, with the exception of the input procedures, which involves capturing the user's set-up and control inputs.

d. Outputs

i. The Mitsuhashi Probe Selection Diagram Window

The Mitsuhashi Probe Selection Diagram (MPSD), FIG. 4, is a key feature of the invention as it is a unique way of visualizing the results of the program's calculations. It is a graphic display of all of the hybridizations of probes for the target mRNA with

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all sequences in the preparation. In other words, given a sequence database and a target mRNA, the MPSD graphically displays all of the candidate probes and their hybridization strengths with all sequences from the sequence database. The MPSD allows the user to see visually the number of false hybridizations at various temperatures for all candidate probes, and the sources of these false hybridizations (with a loci and sequence comparison).

For each melting temperature Tm of interest, a graphical representation of the number of hybridizations for each probe is displayed. In the preferred embodiment, this representation is color coded. In this implementation of the invention, the color red 123 identifies the highest melting temperature Tm and the color blue 124 identifies the lowest melting temperature Tm. Each mismatch results in a reduction in Tm. Tm is also a function of probe length and percent content of GC bases. Within the window, the cursor 125 shape is changed from a vertical line bisecting the screen to a small rectangle when the user selects a particular probe. The current probe is defined to be that probe under the cursor position (whether it be a line or a rectangle) in the MPSD window. More detailed information about the current probe is given in the ProbeInfo and MatchInfo windows, discussed below. Clicking the mouse 2 once at the cursor 125 selects the current probe. Clicking the mouse 2 a second time deselects the current probe. Moving the cursor across the screen causes the display to change to reflect the candidate probe under the current cursor position.

The x-axis 110 of the MPSD, FIG. 4, shows the candidate probes' starting positions along the given mRNA sequence. The user may "slide" the display to the left or right in order to display other probe starting positions. The y-axis 115 of the MPSD displays the probe specificity, which is calculated by the program.

The menu options 116, 117, 118, 119, and 120 available to the user while in the MPSD, FIG. 4, are displayed along a menu bar at the top of the screen. The user can click the mouse 2 on the preferred option to briefly display the option choices, or can click and hold the mouse button on the option to allow an option to be selected. The user may also type a combination of keystrokes in order to display an option in accordance with well-known computer desk top interface operations. This combination usually involves holding down the ALT key while pressing the key representing the first letter of the desired option (i.e, F, P, M, E or H).

The File option 116 allows the user to specify input files and databases. The Preparation option 117 allows the user to create a preparation file summarizing the

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sequence database. The Models option 118 allows the user to specify the hybridization model (i.e., H-Site or Mismatch) and its parameters. The Experiment option 119 and the Help option 120 are not available in the current implementation of this invention. These options are part of the original Main Dialog Window, FIG. 2.

Areas on the graphical display of the MPSD, FIG. 4, where the hybridizations for the optimal probes are displayed are lowest and most similar, such as shown at 121, indicate that the particular sequence displayed is common to all sequences. Areas on the graphical display of the MPSD where the hybridizations for the optimal probes are displayed are highest and most dissimilar, such as shown at 122, indicate that the particular sequence displayed is extremely specific to that particular gene fragment. The high points on the MPSD show many loci in the database, to which the candidate probe will hybridize (i.e., many false hybridizations). The low points show few hybridizations, at least relative to the given database. In other words, the sequence shown at 121 would reflect a probe common to all of the gene fragments tested, such that this probe could be used to detect each of these genes. The sequence shown at 122 would reflect a probe specific to the particular gene fragment, such that this probe could be used to detect this particular gene and no others.

ii. The ProbeInfo and MatchInfo Window

The combined ProbeInfo and MatchInfo Window, FIG. 5, displays detailed information about the current candidate probe. The upper portion of the window is the ProbeInfo window, and the lower portion is the MatchInfo window. The ProbeInfo window portion displays the following types of information: the target locus (i.e., the mRNA, cDNA, or DNA from which the user is looking for probes) is displayed at 131, while the preparation used for hybridizations is displayed at 132. In the example shown in FIG. 5, the target locus 131 is the file named HUMBJUNX.CDS, which is shown as being located on drive F in the subdirectory MILAN. The preparation 132 is shown as being the file designated JUNMIX.PRP, which is also shown as being located on drive F in the subdirectory MILAN. The JUNMIX.PRP preparation in this example is a mixture of human and mouse jun loci.

The current and optimal probe's starting position is shown at 135. The current candidate oligonucleotide probe is defined at 136, and is listed at 137 as having a length of 21 bases. The melting temperature for the probe 136 as hybridized with the targets is shown in column 140. The melting temperature for the optimal probe is given as 61.7 degrees C at 138. The ProbeInfo Window FIG. 5 also displays hairpin characteristics

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of the probe at 139. In the example shown, the ProbeInfo Window shows that there are four (4) base pairs involved in the worst hairpin, and that the worst hairpin has a length of one (1) (see FIG. 5, at 139).

The MatchInfo Window portion displays a list of hybridizations between the current probe and species within the preparation file, including hybridization loci and hybridization temperatures. The hybridizations are listed in descending order by melting temperature. The display shows the locus with which the hybridization occurs, the position within the locus, and the hybridization sequence.

In the MatchInfo window portion, the candidate probe 136 is shown at 150 as hybridizing completely with a high binding strength. This is because the target DNA is itself represented in the database in this case, so the candidate probe is seen at 150 to hybridize with itself (a perfect hybridization). The locus of each hybridization from the preparation 132 are displayed in column 141, while the starting position of each hybridization is given in column 142. The calculated hybridizations are shown at 145.

iii. The ProbesEdit Window

The ProbesEdit Window, FIG. 6, is a text editing window provided for convenient editing and annotation of the invention's text file output. It is also used to accumulate probes selected from the MPSD, FIG. 4, by mouse 2 clicks. Standard text editing capabilities are available within the ProbesEdit Window. The user may accumulate selected probes in this window (see 155 for an example) and then save them to a file (which will bear the name of the preparation sequence with the file extension of "prb" 156, or may be another file name selected by the user). A sample of this file is shown in FIG. 6A.

iv. Miscellaneous Output

The present embodiment of this invention also creates two output files, currently named "test.out" and "test1.out", depending upon which model the user has selected. The first file, "test.out", is created with both the Mismatch Model and the H-Site Model. This file is a textual representation of the Mitsuhashi Probe Selection Diagram (MPSD). It breaks the probe sequence down by position, length, delta Tm, screensN, and the actual probe sequence (i.e., nucleotides). An example of this file created by the Mismatch Model is shown in FIG. 20, and example created by the H-Site Model is shown in FIG. 24A. The second file, "test1.out", is created only by the H-Site Model. This file is a textual representation of the ProbeInfo and MatchInfo window that captures all hybridizations, along with their locus, starting position, melting temperature,

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and possible other hybridizations. A partial example of this file is shown in FIG. 24B (10 pages out of a total of 190 pages created by the H-Site Model).

2. Description of the Mismatch Model Program

a. Overview

In this invention, one of the hybridization strength models is termed the Mismatch Model (see FIG. 2 for selection of this model). The basic operation of this model involves the techniques of hashing and continuous seed filtration, as defined earlier and described in more detail below. The essence of the Mismatch Model is a fast process for doing exact and inexact matching between DNA and mRNA sequences to support the Mitsuhashi Probe Selection Diagram (MPSD). There are a number of modules in the present implementation of the Mismatch Model contained in this invention, the most significant of which are shown in the flow chart in FIG. 7 and in more detail in FIGS. 8 through 18. The main k_diff module shown in the flow chart in FIG. 8 is a structured program that provides overall control of the Mismatch Model, calling various submodules that perform different functions.

b. Inputs

The user-selected input variables for this model are minimum probe length 26 (which is generally from 18 to 30) and maximum number of mismatches 27 (which generally is from 1 to 5). These inputs are entered by the user in the Main Dialog Window, FIG. 2C.

c. **Processing**

i. k diff Program

Some terms of art need to be defined before the processing performed by this module can be explained. A hash table basically is an array or table of data. A linked list is a classical data structure which is a chain of linked entries and involves pointers to other entry structures. Entries in a linked list do not have to be stored sequentially in memory, as is the case with elements contained in an array. Usually there is a pointer to the list associated with the list, which is often initially set to point to the start of the list. A pointer to a list is useful for sequencing through the entries in the list. A null pointer (i.e., a pointer with a value of zero) is used to mark the end of the list.

As the flow charts in FIGS. 7 and 8 illustrate, the general process steps and implemented functions of this model can be outlined as follows:

Step 1: First, create a hash table and linked list from the query (FIG. 7, hashing module 222).

Step 2: Next, while there are still GenBank entries available for searching (FIG. 7, assembly module 230):

Step 2a: Read the current GenBank entry (record) sequence of user-specified length (FIG. 7, seqload module 232), or read the current sequence (record) from the file selected by the user (FIG. 7, read1 module 234).

Step 2b: For the current sequence for each position of the sequence from the first position (or nucleotide) to the last position (or nucleotide) (incrementing the position number once each iteration of the loop) (FIG. 7, q_colour module 242),

Step 2c: set the variable dna_hash equal to the hash of the current position of the current sequence (FIG. 7, q_colour module 242). Step 2d: While not at the end of the linked list for dna_hash (FIG. 7, q_colour module 242),

Step 2e: set the query_pos equal to the current position of dna_hash in the linked list (FIG. 7, q_colour module 242) and

Step 2f: Extend the hit with the coordinates (query_pos, dna_pos) (FIG. 7, hit_ext module 244),

Step 2g: If there exists a k_mismatch in the current extended hit (FIG. 7, colour module 246), then

Step 2h: print the current hit (FIG. 7, q_colour module 242), and repeat from Step 2.

As this illustrates, there are three (3) basic looping or iteration processes with functions being performed based on variables such as whether the GenBank section end has been reached (the first "WHILE" loop, Step 2), whether the end of the current DNA entry has been reached (the "FOR" loop, Step 2b), and whether the end of the dna_hash linked list has been reached (the second "WHILE" loop, Step 2d). A "hit" will only be printed if there are k mismatches in the current extended hit.

FIGS. 8 through 18 illustrate the functions of each of the modules of the present embodiment of this invention, all of which were generalized and summarized in the description above. FIG. 8, which outlines the main "k_diff" module, shows that this

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module is primarily a program organization and direction module, in addition to performing routine "housekeeping" functions, such as defining the variables and hash tables 251, checking if the user-selected gene sequence file is open 252, extracting needed identification information from the GenBank 253, and ensuring valid user input 254. This module also performs a one-time allocation of memory for the gene sequences, and allocates memory for hit information, hashing, hybridization and frequency length profiles and output displays, 255 & 256. The "k_diff" module also initializes or "zeros out" the hashing table, the linked hashing list and the various other variables 257 in preparation for the hashing function. In addition, this module forms the hash tables 258 and extracts a sequence and finds the sequence length 259.

One of the most important functions performed by the "k_diff" module is to define the seed (or kernel or k_tuple) size. This is done by setting the variable k_tuple equal to (min_probe_length - max_mismatch_#)/(max_mismatch+# + 1) FIG. 8 at 265. Next, if the remainder of the aforementioned process is not equal to zero 266, then the value of the variable k_tuple is incremented by one 267. The resulting value is the size of the seed. The module then reads the query 268 and copies the LOCUS name 269 for identification purposes (a definition of the term locus is given earlier in the specification).

The "k_diff" module FIG. 8 also calls the "assembly" module 260, writes the results to a file 261a, plots the results 261b (discussed below), calculates the hairpin characteristics 262 (i.e., the number of base pairs and the length of the worst hairpin) and the melting temperature (Tm) for each candidate probe 263, and saves the results to a file 264.

The screen graphs are plotted 261b by converting the result values to pixels, filing a pixel array and performing a binary search into the pixel array. Next, given the number of pixels per probe position and which function is of interest to the user (i.e., the three mismatch match numbers), the program interpolates the values at the value of (pixelsPerPositionN-1) and computes the array of pixel values for drawing the graph. These values are then plotted on the MPSD.

The "hashing" module, FIG. 9, performs hashing of the query. In other words, it creates the hash table and linked list of query positions with the same hash. The variable has_table[i] equals the position of the first occurrence of hash i in the query. If i does not appear in the query, hash_table[i] is set to zero.

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The "tran" module, FIG. 10, is called by the "hashing" module 271, and performs the hashing of the sequence of k_tuple (kernel or seed) size. If the k_tuple exists (i.e., its length is greater than zero), the variable uns is set equal to uns*ALF+p 291. The variable p represents the digit returned by the "let_dig" module FIG. 11 that represents the nucleotide being examined. ALF is a constant that is set by the program in this implementation to equal four. The query pointer is then incremented, while the size of k_tuple (the seed) is decremented 292. This process is repeated until the sequence of k_tuple has been entirely hashed. Then the "tran" module returns the variable current_hash 293 to the "hashing" module FIG. 9.

The "let_dig" module, FIG. 11, is called by the "tran" module 291, and transforms the nucleotides represented as the characters "A", "T", "U", "G" and "C" in the GenBank and the user's query into numeric digits for easier processing by the program. This module transforms "a" and "A" into "0"301, "t", "T", "u" and "U" into "1"302, "g" and "G" into "2"303, and "c" and "C" into "3"305. If the character to be transformed does not match any one of those listed above, the module returns "-1" 305. The "hashing" module, FIG. 9, then calls the "update" module 272, FIG. 12, which updates the hash with a sliding window (i.e., it forms a new hash after shifting the old hash by "1"). The remainder of old_hash divided by power_1 is calculated 311 (a modulus operation), the remainder is multiplied by ALF 312 (i.e., four), and then the digit representing the nucleotide is added to the result 313. The "update" module then returns the result 314 to the "hashing" module FIG. 9.

If the current hash has already occurred in the query, the program searches for the end of the linked list for the current hash 273 and marks the end of the linked list for the current hash 274. If the current hash has not already occurred in the query, the program puts the hash into the hash table 275. The resulting hash table and linked list are then returned to the "k diff" module, FIG. 8 at 258.

The "assembly" module, FIG. 13, extracts sequences from the GenBank and performs hit locating and extending functions. This module is called by the "k_diff" module FIG. 8 at 260 if the user has chosen to use the database to locate matches. The output from the "assembly" module (FIG. 13) tells the user that the section of the database searched contains E number of entries 321 of S summary length 322 with H number of hits 323. Further, the program tells the user that the number of considered l-tuples equals T 324. The entry head line is also printed 326.

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The "seqload" module, FIG. 14, is called by the "k_diff" module FIG. 8 at 259 once the query hash table and linked list have been formed by the "hashing" module FIG. 9. The "seqload" module FIG. 14 checks to see if the end of the GenBank file has been reached 327, and, if not, searches until a record is found with LOCUS in the head-line 328. Next, the LOCUS name is extracted 329 for identification purposes, and the program searches for the ORIGIN field in the record 330.

The program then extracts the current sequence 331 from the GenBank and performs two passes on each sequence. The first is to determine the sequence length 332 and allocate memory for each sequence 333, and the second pass is to read the sequence into the allocated memory 334. Since the sequences being extracted can contain either DNA nucleotides or protein nucleotides, the "seqload" module can recognize the characters "A","T","U","G",and "C". The bases "A","T","G" and "C" are used in DNA sequences, while the bases "A","U", "G" and "C" are used in RNA and mRNA sequences. The extracted sequence is then positioned according to the type of nucleotides contained in the sequence 335, and the process is repeated. Once the end of the sequence has been reached, the "seqload" module returns the sequence length 336 to the "k_diff" module FIG. 8.

If the user has chosen to use one or more files to locate matches, rather than the database, the "read1" module, FIG. 15, rather than the "seqload" module FIG. 14, is called by the "k_diff"module FIG. 8. The "read1" module, FIG. 15, reads the sequence from the user specified query file 341 and allocates memory 342. This module also determines the query length 343, extracts sequence identification information 344, determines the sequence length 345, transforms each nucleotide into a digit 346 by calling the "let_dig" module FIG. 11, creates the query hash table 347 by calling the "dig_let: module FIG. 16, and closes the file 348 once everything has been read in.

First, the "read1" module FIG. 15 allocates space for the query 342. To do this, the "ckalloc" module, FIG. 15 at 342, is called. This module allocates space and checks whether this allocation is successful (i.e., is there enough memory or has the program run out of memory). After allocating space, the "read1" module FIG. 15 opens the user-specified file 349 (the "ckopen" module, FIG. 15 at 349, is called to ensure that the query file can be successfully opened 349), determines the query length 343, locates a record with LOCUS in the head-line and extracts the LOCUS name 344 for identification purposes, locates the ORIGIN field in the record and then reads the query sequence from the file 341. Next, the sequence length is determined 345, memory is

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allocated for the sequence 342, and the sequence is read into the query file 350. If the string has previously been found, processing is returned to 344. If not, then each character in the query file is read into memory 350.

The characters are transformed into digits 346 using the "let_dig" module, FIG. 11, until a valid digit has been found, and then the hash table containing the query is set up 347 using the module "dig_let", FIG. 16, which transforms the digits into nucleotides represented by the characters "A"371, "T"371, "G"373, "C"374, and "X"375 as a default. If the end of the file has not been reached, processing is returned to 344. If it has, the file is closed 348 and the query is then returned to the "read1" module FIG. 15 at 347.

The "q_colour" module, FIG. 17 (FIG. 13 at 325), is called by the "assembly" module FIG. 13 after the current sequence has been extracted from the GenBank. The "q_colour" module FIG. 17 performs the heart of the Mismatch Model process in that it performs the comparison between the query and the database or file sequences. If the module finds that there exists a long (i.e., greater than the min_hit_length) extended hit, it returns a "1"to the "assembly" module FIG. 14. Otherwise, the "q_colour" module, FIG. 17, returns a "0".

In the "q_colour" module, FIG. 17, all DNA positions are analyzed in the following manner. First, the entire DNA sequence is analyzed 391 to see whether each position is equal to zero 392 (i.e., whether it is empty or the sequence is finished). If it is not equal to zero 393, the "q_colour" module FIG. 20 calls the "tran" module, FIG. 10 described above, which performs the hashing of k_tuples. The "tran" module FIG. 10 calls other modules which transform the nucleotides represented by characters into digits for easier processing by the program and then updates the hash with a sliding window. If the position is equal to zero, the current_hash position is set to new_has after one shift of old_hash 390 by calling the "update" module FIG. 12.

If the nucleotide at the current_hash position is equal to zero, processing is returned to 391. If not, the query position is set equal to (nucleotide at current hash position - 1). Next, the "q_colour" module FIG. 17 looks for the current_hash in the hash table 394. If the current k_tuple does not match the query 395, then the next k_tuple is considered 395, and processing is returned to 391. If the current k_tuple does match the query, then the program checks the hit's (i.e., the match's) vicinity 396 by calling the "hit_ext" module, FIG. 18 to determine if the hit is weak. The inventors have found that if the code for the module "hit_ext" is included within the module "q_colour",

rather than being a separate module utilizing the parameter transfer machinery, 25% of CPU time can be saved.

The "hit_ext" module FIG. 18 determines the current query position in the hit's vicinity 421, determines the current DNA position in the hit's vicinity 422, and creates the list of mismatch positions (i.e., the mismatch_location_ahead 423, the mismatch_location_behind 423 and the kernel match location). If the hit is weak 424, the "hit_ext" module FIG. 18 returns "0" to the "q_colour" module FIG. 17. If the hit has a chance to contain 425, the module returns "1" to the "q_colour" module FIG. 17. A hit has a chance to contain, and is therefore not considered weak, if the mismatch_location_ahead - the mismatch_location_behind is greater than the min_hit_length. If not, it is a short hit and is too weak.

If the "hit_ext" module FIG. 18 tells the "q_colour" module FIG. 17 that the hit was not a weak one, then the "q_colour" module determines whether the current hit is long enough 398 by calling the "colour" module FIG. 19. The "colour" module FIG. 19 performs query_colour modification by the hit data, starting at pos_query and described by mismatch_location_ahead and mismatch_location_behind. After the variables to be used in this module are defined, variable isw_print (which is the switch indicating the hit length) is initialized to zero 430. The cur_length is then set equal to the length of the extending hit 431 (mismatch_location_behind[i] + mismatch_location_ahead[j]-1). Next, if cur_length is greater than or equal to the min_hit_length 432 (i.e., the minimum considered probe size), the hit is considered long and isw_print is set equal to two 433. The value of isw_print is then returned 434 to the "q colour" module FIG. 17.

If the length of the extending hit is longer than the min_hit_length, the hit is considered long 399. Otherwise, the hit is considered short. If the hit is short, nothing more is done to the current hit and the module begins again. If, on the other hand, the hit is considered long 399, the "q_colour" module FIG. 17 prints the current extended hit 400. The current extended hit can be printed in ASCII, printed in a binary file, or printed to a memory file. The "q_colour" module FIG. 17 then repeats until the end of the linked list is reached.

d. Outputs

The output of the k_diff program in the current implementation of this invention may be either a binary file containing the number of extended hits and the k_mismatch hit locations (see FIG. 20), or the output may be kept in memory without writing it to a file. See Section 1(d)(iv) for more detail.

3. Description of the H-Site Model Program

a. Overview

In this invention, the second hybridization strength model is termed the H-Site Model (see FIG. 2 for user selection of this model). One aspect of the H-Site Model uses a generalization of an experimental formula in general usage. The formula used in the H-Site Model is an expression of the fact that melting temperature Tm is a function of both probe length and percent of GC content. This basic formula has been modified in this invention to account for the presence of mismatches. Each percent of mismatch reduces the melting temperature Tm by an average of 1.25 degrees (2 degrees C for an AT mismatch, and 4 degrees C for a GC mismatch).

In addition, this implementation of the invention does some preliminary preprocessing of the GenBank database to sort out and select the cDNA sequences. This is done by locating a keyword (in this case CDS) in each GenBank record. No other programs currently available allow for this combination of functions as far as the inventors are aware.

There are a number of modules in the present embodiment of the H-Site Model contained in this invention. Each step of the processing involved in the H-Site Model is more fully explained below, and is accompanied by detailed flow charts.

b. Inputs

There are two basic user-selected inputs for the H-Site Model (see FIG. 2C): 1) the melting temperature Tm 22 for which probes are being designed (i.e., the melting temperature that corresponds to a particular experiment or condition the user desires to simulate); and 2) the nucleation threshold 23, which is the number of base pairs constituting a nucleation site. The user is also required to select the 1) target species 11 gene sequence(s) (DNA, mRNA or cDNA) for which probes are being designed; 2) the preparation 12 of all sequences with which hybridizations are to be calculated; and 3) the probe output file 13. The preparation file is the most important, as discussed below.

c. Organization of the H-Site Model Program

The current implementation of the H-Site Model program of this invention is distributed between five files containing numerous modules. The main file is designated by the inventors as "ds.cpp" in its uncompiled version. This file provides overall control to the entire invention. It is divided into six sections. Section 0 defines and manipulates global variables. Section 1 controls general variable definition and initialization

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(including the arrays and memory blocks). It also reads and writes buffers for user input selections, and constructs multi buffers.

Section 2 sets up and initializes various "snippet" variables (see section below for a complete definition of the term snippet), converts base pair characters to a representation that is 96 base pairs long and to ASCII base pair strings, and performs other sequence file manipulation such as comparing snippets. This section also reads the sequence format file, reads base pairs, checks for and extracts sequence identification information (such as ORIGIN and LOCUS) and filters out sequences beginning with numbers.

Section 3 involves preparation file manipulation. This section performs the preprocessing on the PRP file discussed above. It also merges and sorts the snippet files, creates a PRP file and sorts it, and outputs the sorted snippets. Next, this section streams through the PRP file.

Section 4 contains the essential code for H-Site Model processing (see FIGS. 21 through 23 for details, discussed below). Streams are set up, and then RIBI comparisons are performed for hybridizations (see file "ribi.cpp" for definitions of RIBI search techniques). Next, probes are generated, binding strength is converted to melting temperature, and hybridizations are calculated and stored (including hybridization strength). Lastly, other H-Site calculations are performed.

Section 5 is concerned with formatting and presenting diagnostic and user file (test.out, test1.out, and test2.out files) output. This section also handles the graphing functions (the MPSD diagram in particular). In addition, this section calculates the hairpin characteristics for the H-Site Model candidate probes.

The second H-Site Model file, designated as "ds.h" defines data variables and structures. Section 1 of this file concerns generic data structures (including memory blocks and arrays, and file inputs and outputs). Section 2 defines the variables and structures used with sequences, probes and hybridizations. Section 3 defines variables and structures concerned with protocols (i.e., function prototypes, graphing, etc.).

The third H-Site Model file, designated as "funcdoc.txt", contains very detailed documentation for this implementation of the H-Site Model program. Numerous variables and structures are also defined. The flow of the program is clearly shown in this file.

The fourth H-Site Model file, designated as "ribi.h" handles the sequence comparisons. The fifth and last H-Site Model file, designated as "ribi.cpp", performs

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internal B-Tree indexing. Definitions of Red-black Internal Binary Index (RIBI) searching are found in this file. Definitions are also included for the concepts keyed set, index, binary tree, internal binary index, paths, and red-black trees. Implementation notes are also included in this file.

d. Processing

Implementation of the H-Site Model in this invention is done in three stages. First, the invention creates the preparation (PRP) file, which contains all relevant information from the sequence database. This is the preprocessing stage discussed above. Next, the target is prepared by the program. Lastly, the invention calculates the MPSD data using the PRP file and target sequence to find probes.

i. Creation of the Preprocessed Preparation File

FIG. 21. Step 1: The program first opens the sequence database for reading into memory 461, 462. Step 2: Next, as sequence base pairs are read in 462, "snippets" are saved to disk 463, along with loci information. A snippet is a fixed-length subsequence of a preparation sequence. The purpose of snippets is to allow the user to examine a small portion of a preparation sequence together with its surrounding base pairs. Snippets in the implementation of this invention are 96 base pairs long (except for snippets near the end or beginning of a sequence, which may have fewer base pairs). The "origin" of the snippet is in position 40. For snippets taken near the beginning of a sequence, some of the initial 40 bases are undefined. For snippets near the end of a sequence, some of the final 55 bases are undefined. Snippets are arranged in the preparation file (PRP) in sorted order (lexicographical order beginning at position 40). In this invention, the term "lexicographical order" means a preselected order, such as alphabetical, numeric or alphanumeric. In order to conserve space, snippets are only taken at every 4th position of the preparation sequence.

Step 3: The snippets are merge sorted 464 to be able to search quickly for sequences which pass the "screen", discussed below. Step 4: The merged file is prepended with identifiers for the sources of the snippets 465. This is done to identify the loci from which hybridizations arise.

ii. Target Preparation

FIG. 22. Step 1: The target sequence file is opened 471 and read into memory 472. For each position in the target mRNA, the probe defined at that starting position is the shortest subsequence starting at that position whose hybridization strength is greater than the user specified melting temperature Tm. Typically, the probes are of

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length 18 to 50. Step 2: Four lists of "screens" are formed 473, 474, 475, each shifted by one base pair 475 to correspond to the fact that snippets are only taken at every four base pairs. A screen is a subsequence of the target mRNA of length equal to the screening threshold specified by the user. The screens are then indexed 476 and sorted in memory 477.

iii. Calculation of the MPSD Data

FIG. 23. Step 3: This step is the heart of the process. Step 3a: The program streams through the following five items in sync, examining them in sequential order: the snippet file and the four lists of screens 481-484. Step 3b: Each snippet is compared to a screen 485. Step 3c: If the snippet does not match, whichever stream is behind is advanced 486 and Step 3b is repeated. If the snippet does match, Step 4 is performed.

Step 4: If a snippet and a matching screen were found in Step 3b 487, the hybridization strength of the binding between the sequence containing the snippet and all of the probes containing the screen is calculated (see Step 5). Double counting is avoided by doing this only for the first matched screen containing the probe. Each pair of bases is examined and assigned a numerical binding strength. An AT pair would be assigned a lower binding strength than a GC pair because AT pairs have a lower melting temperature Tm. The process is explained more fully below at Step 5b.

Step 5: The hybridization strengths between sequence and all the probes containing it are calculated using a dynamic programming process. The process is as follows: Step 5a: Begin at the position of the first probe containing the given screen but not containing any other screens which start at an earlier position and also match the sequence. This is done to avoid double counting. Two running totals are maintained: a) boundStrength, which represents the hybridization strength contribution which would result if the sequence and probe were to match exactly for all base pairs to the right of the current position, and b) unboundStrength, which represents the strength of the maximally binding region. Step 5b: At each new base pair, the variable boundStrength is incremented by 71 if the sequence and probe match and the matched base pair is GC 489, incremented by 30 if the matched base pair is AT 490 (i.e., this number is about 42.25% of the first number 71), and decremented by 74.5 if there is not a match 488 (i.e., this number is about 5% larger than the first number 71). Step 5c: If the current boundStrength exceeds the current unboundStrength 491 (which was originally initialized to zero), a new binding region has been found, and

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unboundStrength is set equal to boundStrength 492. Step 5d: If the current boundStrength is negative, boundStrength is reset to zero 493. Step 5e: If the current position is at the end of a probe, the results (the hybridization strengths) are tallied for that probe. Step 5f: If the current position is at the end of the last probe containing the screen, the process stops.

Step 6: A tally is kept of the number and melting temperature of the matches for each candidate probe, and the location of the best 20 candidates, using a priority queue (reverse order by hybridization strength number) 494. Step 7: A numerical "score" is kept for each preparation sequence by tallying the quantity exp (which can be expressed as Σe^{-Tm}) for each match 495, where Tm is the melting temperature for the "perfect" match, the probe itself. In other words, the probe hybridizes "perfectly" to its target.

Step 8: Hairpins are calculated by first calculating the complementary probe. In other words, the order of the bases in the candidate probe are reversed (CTATAG to GATATC), and complementary base pairs are substituted (A for T, T for A, G for C, and C for G, changing GATATC to CTATAG in the above example). Next, the variable representing the maximum hairpin length for a candidate probe is initialized to zero, as is the variable representing a hairpin's distance. For each offset, the original candidate probe and the complementary probe just created are then aligned with each other and compared. The longest match is then found. If any two matches have the same length, the one with the longest hairpin distance (i.e., the number of base pairs separating the match) is then saved.

Step 9: The preparation sequences are then sorted 496 and displayed in rank order, from best to worst 497. Step 10: The resulting MPSD, which includes <u>all</u> candidate probes, is then displayed on the screen. Step 11: The best 20 matches are also printed or displayed in rank order, as the user requests 497.

e. Outputs

The outputs of the H-Site Model as currently implemented in this invention are fully described in Section 1(d)(iv), above, and illustrated in FIGS. 4 through 6. Samples of the two output files created by the H-Site Model are shown in FIGS. 24A and 24B.

4. <u>Description of the Mitsuhashi Probe Selection Diagram Processing</u>

Once the Mitsuhashi Probe Selection Diagram (MPSD) data has been calculated by the H-Site Model program (see stage three and FIG. 23, discussed above), it is

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necessary to convert this data to pixel format and plot a graph. An overview of this process is shown in FIG. 25. First, the program calculates the output (x,y) ranges 500. Next, these are converted to a logarithmic scale 501. The values are then interpolated 502, and a bitmap is created 503. Lastly, the bitmap is displayed on the screen 504 in MPSD format (discussed above in section 1(e)(i)). A sample MPSD is shown in FIG. 4.

5. Description of the MatchInfo Window Processing

The ProbeInfo and MatchInfo windows are discussed in great detail in Section 1(e)(ii), and a sample of these windows is shown in FIG. 5. An overview of the processing involved in creating the MatchInfo portion of the window is given in the flow chart in FIG. 26. First, as the user moves the MPSD cursor 520 (seen as a vertical line bisecting the MPSD window), the program updates the position of the candidate probe shown under that cursor position 521. Next, based upon the candidate probe's position, the program updates the sequence 522 and hairpin information 523 for that probe. This updated information is then displayed in an updated match list 524, shown in the MatchInfo window.

The above described embodiments of the present invention are merely descriptive of its principles and are not to be considered limiting. The scope of the present invention instead shall be determined from the scope of the following claims including their equivalents.

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WHAT IS CLAIMED IS:

1. A programmed computer system for designing optimal oligonucleotide sequences for use with a gene sequence data source comprising:

first input means for introducing user-selected gene sequence into the computer system;

memory means for storing user-selected gene sequence;

means for accessing gene sequence data from said gene sequence data source;

means for performing exact and inexact match modeling between gene sequences;

means for performing hybridization strength modeling on gene sequences; means for selecting either of said modeling means; and

means for presenting the results of said modeling to present candidate oligonucleotide sequences.

2. A programmed computer system in accordance with Claim 1 wherein said means for performing exact and inexact match modeling utilizes said accessing means to introduce a user-selected set of gene sequence data and a user-selected set of target gene sequence data from said gene sequence data source into the computer system and said memory means to store said gene sequence data and said target gene sequence data and wherein said means for performing exact and inexact match modeling includes:

means for determining a minimum sequence length;

means for creating a look-up hash table and linked list in memory for each gene sequence in said gene sequence data and each of said target gene sequences;

means for calculating the minimum length of any matching gene subsequence of said gene sequence data and said target gene sequence data;

means for comparing each base pair character in each said target sequence stored in a hash table in memory to each base pair character of said gene sequence stored in a hash table in memory;

means for finding a matching seed by determining if the said comparison results in a matching gene subsequence of length equal to said calculated minimum length;

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means for comparing base pair characters behind and ahead of said seed to determine if there exists an extended match of a subsequence of base pair characters of length greater than the calculated minimum length, resulting in a current hit sequence;

means for calculating whether said current hit sequence is longer than said minimum sequence length, resulting in a current candidate oligonucleotide sequence;

means for storing said current candidate oligonucleotide sequence; and wherein said presenting means provides said current candidate oligonucleotide sequence to the user.

3. A programmed computer system in accordance with Claim 2 wherein said computer system includes:

means for calculating the melting temperature for each candidate oligonucleotide sequence;

means for tracking the number and melting temperature of the matches for each candidate oligonucleotide sequence;

means for tracking the location of a set number of the best candidate oligonucleotide sequences; and

wherein said presenting means is operative to present said additional results to the user; and

wherein said presenting means provides said melting temperature to the user.

4. A programmed computer system in accordance with Claim 2 wherein said computer system includes:

means for determining the length of sequences from said target gene sequence data.

5. A programmed computer system in accordance with Claim 2 wherein said computer system includes:

means for determining the length of sequences from said set of gene sequence data.

6. A programmed computer system in accordance with Claim 2 wherein said computer system includes:

means for copying the LOCUS name for each said gene sequence into said memory means; and

means for linking said LOCUS name with each said gene sequence.

- 7. A programmed computer system in accordance with Claim 2 wherein said means for performing exact and inexact match modeling utilizes said accessing means to introduce a user-selected minimum sequence length from said gene sequence data source into the computer system and said memory means to store said minimum sequence length.
- 8. A programmed computer system in accordance with Claim 2 wherein said computer system includes:

means for calculating the melting temperature for each candidate oligonucleotide sequence;

means for tracking the number and melting temperature of the matches for each candidate oligonucleotide sequence;

means for tracking the location of a set number of the best candidate oligonucleotide sequences employing a priority queue by sorting said candidate oligonucleotide sequences in reverse order and sorting said candidate oligonucleotide sequences by hybridization strength;

wherein said presenting means is operative to present said additional results to the user; and

wherein said presenting means provides said melting temperature to the user.

9. A programmed computer system in accordance with Claim 2 wherein said first input means in operative to introduce a user-selected maximum number of mismatches and a user-selected minimum candidate oligonucleotide sequence length into the computer system, and wherein said means for calculating the minimum length of any matching gene subsequence of said gene sequence data and said target gene sequence data comprises the steps of:

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means for subtracting said maximum number of mismatches from said minimum candidate oligonucleotide sequence length to give a first result;

means for dividing said first result by said maximum number of mismatches plus one to give a second result;

means for incrementing said second result by one if the remainder is not equal to zero to give a third result; and

means for truncating said third result to an integer.

10. A programmed computer system in accordance with Claim 9 wherein said means for calculating the hairpin characteristics of said candidate oligonucleotide sequence comprises the steps of:

calculating a complementary sequence to the candidate oligonucleotide sequence by reversing the base pair order of the candidate oligonucleotide sequence and substituting complementary base pairs;

comparing each character of said original candidate oligonucleotide sequence and said complementary sequence;

finding the longest match between said original candidate oligonucleotide sequence and said complementary sequence; and

saving the match with the longest hairpin distance if any two matches have the same length;

means for storing hairpin characteristics; and wherein said presenting means provides said hairpin characteristics to the user.

- 11. A programmed computer system in accordance with Claim 2 wherein said computer system includes a means for calculating the hairpin characteristics of said candidate oligonucleotide sequence.
- 12. A programmed computer system in accordance with Claim 2 wherein said means for preprocessing said set of target gene sequence data and said set of gene sequence data comprises the steps of:

searching for sequences without introns in said target gene sequences and said gene sequences;

extracting target gene sequences and gene sequences that do not contain introns; and

storing said extracted target gene sequences and gene sequences in memory.

13. A programmed computer system in accordance with Claim 1 wherein said means for performing hybridization strength modeling utilizes said first input means to introduce a user-selected screening threshold into the computer system and said accessing means to introduce a user-selected set of gene sequence data and a user-selected set of target gene sequence data from said gene sequence data source into the computer system, and said memory means to store said gene sequence data, said target gene sequence data and said screening threshold and wherein said means for performing hybridization strength modeling comprises:

means for preprocessing said target gene sequence data and said gene sequence data by selecting only those sequences without introns;

means for forming a preparation file of gene sequence fragments by cutting said target gene sequences into fixed length target gene subsequences and sorting said subsequences in lexicographical order;

means for merge sorting said gene sequences;

means for forming multiple lists of screens by forming lists of subsequences of the preparation file of length equal to said screening threshold;

means for indexing, sorting and storing said screens in said memory means;
means for sequentially comparing said preparation file gene sequences with
each of said screens to design candidate oligonucleotide sequences;

means for calculating the hybridization strengths between a gene sequence and all candidate oligonucleotide sequences containing that gene sequence by accounting for Guanine-Cytosine (GC) and Adenine-Thymine (AT) base pair content of the gene sequence and the number of mismatches between said preparation file sequences and a said screen when said comparison results in a match;

means for preparing the candidate oligonucleotide sequence and hybridization strength for presentation to the user; and

wherein said presenting means provides the candidate oligonucleotide sequence and hybridization strength to the user.

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14. A programmed computer system in accordance with Claim 13 wherein said computer system includes:

means for calculating the melting temperature for each candidate oligonucleotide sequence;

means for tracking the number and melting temperature of the matches for each candidate oligonucleotide sequence;

means for tracking the location of a set number of the best candidate oligonucleotide sequences;

means for preparing the melting temperature for presentation to the user; and

wherein said presenting means provides the melting temperature to the user.

15. A programmed computer system in accordance with Claim 14 wherein said means for calculating said candidate oligonucleotide sequence's melting temperature comprises:

solving the formula Tm = 81.5 - 16.6(log[Na]) - .63% (formamide) + ((.41 (%(G + C)) - 600)/N), wherein log[Na] is the sodium concentration, %(G + C) is the fraction of matched base pairs which are G-C complementary, N is the sequence length and wherein the number of mismatches is equal to zero.

16. A programmed computer system in accordance with Claim 15 wherein said computer system includes:

means for reducing a candidate oligonucleotide probe's calculated melting temperature by a certain amount for each percent of mismatch between the candidate oligonucleotide sequence and a user-selected target gene sequence based upon the assumption that there are an equal number of GC and AT base pair mismatches.

17. A programmed computer system in accordance with Claim 16 wherein said means for reducing a candidate oligonucleotide sequence's calculated melting temperature comprises the steps of:

reducing said calculated melting temperature by 2 degrees Celsius if an AT mismatch exists; and

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reducing said calculated melting temperature by 4 degrees Celsius if a GC mismatch exists.

18. A programmed computer system in accordance with Claim 13 wherein said computer system includes:

means for assigning a numerical score to each said gene sequence; and means for sorting said gene sequences in accordance with said numerical score.

19. A programmed computer system in accordance with Claim 13 wherein said means for performing hybridization strength modeling utilizes said accessing means for copying the LOCUS name for each said gene sequence into said memory means, and said memory means; and

means for prepending said gene sequence with said LOCUS name.

- 20. A programmed computer system in accordance with Claim 13 wherein four lists of screens are formed by said list forming means.
- 21. A programmed computer system in accordance with Claim 13 wherein said computer system includes a means of shifting each screen by at least one base pair as it is formed by said list forming means.
- 22. A programmed computer system in accordance with Claim 13 wherein said computer system includes:

means for calculating the melting temperature for each candidate oligonucleotide sequence;

means for tracking the number and melting temperature of the matches for each candidate oligonucleotide sequence;

means for tracking the location of a set number of the best candidate oligonucleotide sequences employing a priority queue by sorting said candidate oligonucleotide sequences in reverse order and sorting said candidate oligonucleotide sequences by hybridization strength;

means for preparing the melting temperature for presentation to the user; and

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wherein said presenting means provides the melting temperature to the user.

23. A programmed computer system in accordance with Claim 13 wherein said computer system includes:

means for assigning a numerical score to each said gene sequence by tallying the quantity "exp" where "exp" = $\Sigma e^{-\tau m}$ and wherein Tm is the melting temperature for the said gene sequence; and

means for sorting said gene sequences in accordance with said numerical score.

24. A programmed computer system in accordance with Claim 13 wherein said means for calculating the hybridization strengths between a gene sequence and all candidate oligonucleotide sequences containing that gene sequence comprises the steps of:

accessing gene sequence data from said gene sequence data source; comparing base pairs of a first gene sequence and a second gene sequence to determine if a match exists;

incrementing said first gene sequence's bound strength by some first number if a base pair character in said first gene sequence and said second gene sequence match and the matched base pair is equal to a combination of the bases Guanine (G) and Cytosine (C);

incrementing said first gene sequence's bound strength by some second number if a base pair character in said first gene sequence and said second gene sequence match and the matched base pair is equal to a combination of the bases Adenine (A) and Thymine (T);

decrementing said first gene sequence's bound strength by a third number if there is no match in base pairs between said first gene sequence and said second gene sequence;

comparing said first gene sequence's bound strength to said first gene sequence's unbound strength;

setting said first gene sequence's unbound strength equal to its bound strength if said first gene sequence's bound strength is greater than said first gene sequence's unbound strength; and

resetting said first gene sequence's bound strength to zero if said first gene sequence's unbound strength is less than zero.

- 25. A programmed computer system in accordance with Claim 24 wherein said first and second numbers are greater than zero.
- 26. A programmed computer system in accordance with Claim 24 wherein said second number is in the order of 42% of said first number.
- 27. A programmed computer system in accordance with Claim 24 wherein said third number is in the order of 5% larger than said first number.
- 28. A programmed computer system in accordance with Claim 13 wherein said computer system includes a means for calculating the hairpin characteristics of said candidate oligonucleotide sequence;

means for preparing the hairpin characteristics for presentation to the user; and

wherein said presenting means provides the hairpin characteristics to the user.

29. A programmed computer system in accordance with Claim 28 wherein said means for calculating the hairpin characteristics of said candidate oligonucleotide sequence comprises the steps of:

calculating a complementary sequence to the candidate oligonucleotide sequence by reversing the base pair order of the candidate oligonucleotide sequence and substituting complementary base pairs;

comparing each character of said original candidate oligonucleotide sequence and said complementary sequence;

finding the longest match between said original candidate oligonucleotide sequence and said complementary sequence; and

saving the match with the longest hairpin distance if any two matches have the same length;

means for preparing the hairpin characteristics for presentation to the user; and

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wherein said presenting means provides the hairpin characteristics to the user.

30. A programmed computer system in accordance with Claim 13 wherein said fixed-length subsequences are calculated by a method comprising the steps of:

locating the origin of said subsequence in a set position of said target gene sequence in said preparation file;

cutting a subsequence that is a fixed-length long every preselected number of positions of said target gene sequence in said preparation file; and

sorting said subsequences in said preparation file in lexicographical order beginning at a set position.

- 31. A programmed computer system in accordance with Claim 30 wherein the origin of said subsequence is located at position 40 of said target sequence in said preparation file.
- 32. A programmed computer system in accordance with Claim 13 wherein said fixed-length subsequences are calculated by a method comprising the steps of:

locating the origin of said subsequence in the 40th position of said target gene sequence in said preparation file;

cutting a subsequence that is 96 base pairs long of said target gene sequence in said preparation file; and

sorting said subsequences in said preparation file in lexicographical order beginning at a set position.

- 33. A programmed computer system in accordance with Claim 13 wherein said computer system includes means for prepending said preparation file subsequences with identifiers for the sources of each subsequence.
- 34. A programmed computer system in accordance with Claim 1 wherein said presenting means to provide the results of said matching and modeling to display candidate oligonucleotide sequences includes means for displaying in multiple dimensions the gene sequences which result from the comparisons and calculations characterized in that said display format exhibits

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the starting position of each candidate oligonucleotide sequence in one dimension;

the specificity of a candidate oligonucleotide sequence's hybridization with the target gene sequence in a second dimension; and

superimposed melting temperatures of gene sequences in contrasting presentations in at least an apparent third dimension.

35. A programmed computer system in accordance with Claim 34 wherein said display further includes a cursor moveable along one dimension of said display that selects a position for an expansion of data representing the homology between the candidate oligonucleotide sequences and said gene sequence data; and

wherein said display is operative to display in alphanumeric form the homology between the candidate oligonucleotide sequences and said gene sequence data.

36. A programmed computer system in accordance with Claim 34 wherein said display is further operative to provide an expansion of data including presenting

false hybridizations at various melting temperatures for all candidate oligonucleotide sequences;

the location of each false hybridization;

a candidate oligonucleotide sequence's starting position; and

hairpin characteristics of each candidate oligonucleotide sequence.

- 37. A programmed computer system in accordance with Claim 34 wherein said display format data is outputted to a printing means.
- 38. A programmed computer system in accordance with Claim 34 wherein said display format data is saved to a data file.
- 39. A programmed computer system in accordance with Claim 34 wherein said display format data is exported to another computer system.

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40. A programmed computer system in accordance with Claim 34 wherein said display further includes a cursor moveable along one dimension of said display that selects a position for an expansion of data representing the homology between the candidate oligonucleotide sequences and said gene sequence data; and

wherein said moveable cursor may be positioned by the user to select and save particular candidate oligonucleotide sequence information; and

wherein said display is operative to display in alphanumeric form the homology between the candidate oligonucleotide sequences and said gene sequence data.

- 41. A programmed computer system in accordance with Claim 40 wherein said method of selecting and saving particular candidate oligonucleotide sequence information comprises capturing candidate oligonucleotide sequence information at the user-selected point and storing said information in said memory means.
- 42. A programmed computer system in accordance with Claim 41 wherein said user-selected candidate oligonucleotide sequence information is exported to another computer system.
- 43. A programmed computer system in accordance with Claim 34 wherein said means for displaying comprises the steps of:

calculating display output ranges; converting said output ranges to a logarithmic scale; interpolating said converted values; creating a bitmap of said interpolations; and displaying said bitmap on a display device.

44. A programmed computer system in accordance with Claim 34 wherein said means for displaying comprises the steps of:

converting said result values to pixels;

filling a pixel array with said pixels;

performing a binary search into said pixel array;

determining the number of pixels per candidate oligonucleotide sequence to be displayed;

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interpolating said pixels at the value of pixels per position minus one; computing an array of said pixel array; and plotting the results on a display device.

45. A programmed computer system in accordance with Claim 1 wherein said means for performing exact and inexact match modeling utilizes said accessing means to introduce a user-selected set of gene sequence data and a user-selected set of target gene sequence data from said gene sequence data source into the computer system and said memory means to store said gene sequence data and said target gene sequence data and wherein said means for performing exact and inexact match modeling includes:

means for determining a minimum sequence length;

means for creating a look-up hash table and linked list in memory for each gene sequence in said gene sequence data and each of said target gene sequences;

means for calculating the minimum length of any matching gene subsequence of said gene sequence data and said target gene sequence data;

means for transforming base characters in each said target sequence and in each said gene sequence into numeric digits;

means for comparing each base pair digit in each said target sequence stored in a hash table in memory to each base pair digit of said gene sequence stored in a hash table in memory;

means for finding a matching seed by determining if the said comparison results in a matching gene subsequence of length equal to said calculated minimum length;

means for comparing base pair digits behind and ahead of said seed to determine if there exists an extended match of a subsequence of base pair digits of length greater than the calculated minimum length, resulting in a current hit sequence;

means for calculating whether said current hit sequence is longer than said minimum sequence length, resulting in a current candidate oligonucleotide sequence;

means for storing said current candidate oligonucleotide sequence; and wherein said presenting means provides said current candidate oligonucleotide sequence to the user.

46. A programmed computer system for designing candidate oligonucleotide sequences for use with a gene sequence data source including:

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first input means for introducing user-selected gene sequence, design, model and presentation criteria and a user-specified sequence length into the computer system;

memory means for storing said gene sequence, design, model and presentation criteria and said sequence length;

means for accessing gene sequence data from said gene sequence data source;

wherein said accessing means is operative to introduce a user-selected set of gene sequence data and a user-selected set of target gene sequence data from said gene sequence data source into the computer system;

wherein said criteria are used for comparison of gene sequence data and target gene sequence data;

means for comparing said gene sequences against said target gene sequences employing said criteria;

means for calculating candidate oligonucleotide sequences of said sequence length that are either common to a pool of user-specified gene sequences or specific to a particular user-specified gene sequence;

means for calculating the homology between the candidate oligonucleotide sequences and said gene sequence data;

means for calculating a candidate oligonucleotide sequence's hairpin characteristics;

means for displaying in multiple dimensions the gene sequences which result from the comparisons and calculations characterized in that said display format exhibits:

the starting position of each candidate oligonucleotide sequence in one dimension;

a candidate oligonucleotide sequence's specificity to the target gene sequence in a second dimension; and

superimposed melting temperatures of gene sequences in contrasting presentations in at least an apparent third dimension;

wherein said display further includes a cursor moveable along one dimension of said display that selects a position for an expansion of data representing

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the homology between the candidate oligonucleotide sequences and said gene sequence data;

wherein said display is operative to display in alphanumeric form the homology between the candidate oligonucleotide sequences and said gene sequence data; and

wherein said display is operative to provide an expansion of data including presenting

false hybridizations at various melting temperatures for all candidate oligonucleotide sequences;

the location of each false hybridization;

a candidate oligonucleotide sequence's starting position; and

hairpin characteristics of each candidate oligonucleotide sequence.

47. A method for designing candidate oligonucleotide sequences by performing exact and inexact match modeling for use with a gene sequence data source comprising the steps of:

introducing user-selected gene sequence into a computer system; accessing gene sequence data from said gene sequence data source; storing user-selected gene sequence in the memory of the computer

accessing the gene sequence source to introduce the user-selected set of gene sequence data and a user-selected set of target gene sequence data from said gene sequence data source into the computer system;

storing said gene sequence data and said target gene sequence data in the memory of the computer system;

determining a minimum sequence length;

system;

creating a look-up hash table and linked list in memory for each gene sequence in said gene sequence data and each of said target gene sequences;

calculating the minimum length of any matching gene subsequence of said gene sequence data and said target gene sequence data;

comparing each base pair character in each said target sequence stored in a hash table in memory to each base pair character of said gene sequence stored in a hash table in memory;

determining a matching seed by determining if the said comparison results in a matching gene subsequence of length equal to said calculated minimum length;

comparing base pair characters behind and ahead of said seed to determine if there exists an extended match of a subsequence of base pair characters of length greater than the calculated minimum length, resulting in a current hit sequence;

calculating whether said current hit sequence is longer than said minimum sequence length, resulting in a current candidate oligonucleotide sequence;

storing said current candidate oligonucleotide sequence in the memory of the computer system; and

presenting a representation of said current candidate oligonucleotide sequence to the user.

48. A method in accordance with Claim 47 wherein said method includes the steps for performing additional calculations for each candidate oligonucleotide probe, said additional calculations comprising:

calculating the melting temperature for each candidate oligonucleotide sequence;

tracking the number and melting temperature of the matches for each candidate oligonucleotide sequence;

tracking the location of a set number of the best candidate oligonucleotide sequences; and

presenting said additional results to the user.

- 49. A method in accordance with Claim 47 wherein said method includes the step of transforming base characters into numeric digits.
- 50. A method in accordance with Claim 47 wherein said method includes the step of determining the length of sequences from said target gene sequence data.
- 51. A method in accordance with Claim 47 wherein said method includes the step of determining the length of sequences from said set of gene sequence data.

52. A method in accordance with Claim 47 wherein said method includes the steps of:

copying the LOCUS name for each said gene sequence into the memory of the computer system; and

linking said LOCUS name with each said gene sequence.

53. A method in accordance with Claim 47 wherein said method includes the steps of:

introducing a user-selected minimum sequence length into the computer system; and

storing said minimum sequence length in the memory of the computer system.

54. A method in accordance with Claim 47 wherein said method includes the steps for performing additional calculations for each candidate oligonucleotide probe, said additional calculations comprising:

calculating the melting temperature for each candidate oligonucleotide sequence;

tracking the number and melting temperature of the matches for each candidate oligonucleotide sequence;

tracking the location of a set number of the best candidate oligonucleotide sequences employing a priority queue by sorting said candidate oligonucleotide sequences in reverse order and sorting said candidate oligonucleotide sequences by hybridization strength; and

presenting said additional results to the user.

55. A method in accordance with Claim 47 wherein said step for calculating the minimum length of any matching gene subsequence comprises:

introducing a user-selected maximum number of mismatches and a user-selected minimum candidate oligonucleotide sequence length into the computer system;

subtracting said maximum number of mismatches from said minimum candidate oligonucleotide sequence length to give a first result;

dividing said first result by said maximum number of mismatches plus one to give a second result;

incrementing said second result by one if the remainder is not equal to zero to give a third result; and

truncating said third result to an integer.

- 56. A method in accordance with Claim 47 wherein said method includes the step of calculating the hairpin characteristics of said candidate oligonucleotide sequence.
- 57. A method in accordance with Claim 47 wherein said method includes the step of calculating the hairpin characteristics of said candidate oligonucleotide sequence comprising:

calculating a complementary sequence to the candidate oligonucleotide sequence by reversing the base pair order of the candidate oligonucleotide sequence and substituting complementary base pairs;

comparing each character of said original candidate oligonucleotide sequence and said complementary sequence;

finding the longest match between said original candidate oligonucleotide sequence and said complementary sequence; and

saving the match with the longest hairpin distance if any two matches have the same length.

58. A method for designing candidate oligonucleotide sequences by performing hybridization strength modeling for use with a gene sequence data source comprising the steps of:

introducing user-selected gene sequence and a user-selected screening threshold into a computer system;

storing user-selected gene sequence and said screening threshold in the memory of the computer system;

accessing the gene sequence source to introduce the user-selected set of gene sequence data and a user-selected set of target gene sequence data from said gene sequence data source into the computer system;

storing said gene sequence data and said target gene sequence data in the memory of the computer system;

preprocessing said target gene sequence data and said gene sequence data by selecting only those sequences without introns;

forming a preparation file of gene sequence fragments by cutting said target gene sequences into fixed length target gene subsequences and sorting said subsequences in lexicographical order;

merge sorting said gene sequences;

forming multiple lists of screens by forming lists of subsequences of the preparation file of length equal to said screening threshold;

indexing and sorting said screens in memory;

storing said screens in the memory of the computer system;

sequentially comparing said preparation file gene sequences with each of said screens to design candidate oligonucleotide sequences;

calculating the hybridization strengths between a gene sequence and all candidate oligonucleotide sequences containing that gene sequence by accounting for Guanine-Cytosine (GC) and Adenine-Thymine (AT) base pair content of the gene sequence and the number of mismatches between said preparation file sequences and a said screen when said comparison results in a match;

preparing the candidate oligonucleotide sequence and hybridization strength for presentation to the user; and

presenting the candidate oligonucleotide sequence and hybridization strength to the user.

59. A method in accordance with Claim 58 wherein said method includes the steps for performing additional calculations for each candidate oligonucleotide probe, said additional calculations comprising:

calculating the melting temperature for each candidate oligonucleotide sequence;

tracking the number and melting temperature of the matches for each candidate oligonucleotide sequence;

tracking the location of a set number of the best candidate oligonucleotide sequences; and

presenting said additional results to the user.

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60. A method in accordance with Claim 58 wherein the step for preparing the candidate oligonucleotide sequence for presenting to the user comprises:

assigning a numerical score to each said gene sequence;

sorting said gene sequences in accordance with said numerical score; and displaying a representation of the resulting candidate oligonucleotide sequence and said gene sequences.

61. A method in accordance with Claim 58 wherein said method includes the steps of:

copying the LOCUS name for each said gene sequence into the memory of the computer system; and

prepending said gene sequence with said LOCUS name.

- 62. A method in accordance with Claim 58 wherein the step for forming lists of screens produces four lists of screens.
- 63. A method in accordance with Claim 58 wherein said method includes a the step of shifting each screen by one base pair as it is formed.
- 64. A method in accordance with Claim 58 wherein said method includes the steps for performing additional calculations for each candidate oligonucleotide probe, said additional calculations comprising:

calculating the melting temperature for each candidate oligonucleotide sequence;

tracking the number and melting temperature of the matches for each candidate oligonucleotide sequence;

tracking the location of a set number of the best candidate oligonucleotide sequences employing a priority queue by sorting said candidate oligonucleotide sequences in reverse order and sorting said candidate oligonucleotide sequences by hybridization strength; and

presenting said additional results to the user.

65. A method in accordance with Claim 58 wherein said method for preparing the results for presenting to the user comprises:

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assigning a numerical score to each said gene sequence by tallying the quantity "exp" where "exp" = Σe^{-Tm} and wherein Tm is the melting temperature for the said gene sequence;

sorting said gene sequences in order of the numerical score; and displaying a representation of the resulting candidate oligonucleotide sequence and said gene sequences.

66. A method in accordance with Claim 58 for use with a gene sequence data source, programmed to determine hybridization strength comprising the steps of:

comparing base pairs of a first gene sequence and a second gene sequence to determine if a match exists;

incrementing said first gene sequence's bound strength by some first number if a base pair character in said first gene sequence and said second gene sequence match and the matched base pair is equal to a combination of the bases Guanine (G) and Cytosine (C);

incrementing said first gene sequence's bound strength by some second number if a base pair character in said first gene sequence and said second gene sequence match and the matched base pair is equal to a combination of the bases Adenine (A) and Thymine (T);

decrementing said first gene sequence's bound strength by a third number if there is no match in base pairs between said first gene sequence and said second gene sequence;

comparing said first gene sequence's bound strength to said first gene sequence's unbound strength;

setting said first gene sequence's unbound strength equal to its bound strength if said first gene sequence's bound strength is greater than said first gene sequence's unbound strength; and

resetting said first gene sequence's bound strength to zero if said first gene sequence's unbound strength is less than zero.

67. A method in accordance with Claim 66 wherein said first and second numbers are greater than zero.

- 68. A method in accordance with Claim 66 wherein said second number is in the order of 42% of said first number.
- 69. A method in accordance with Claim 66 wherein said second number is in the order of 5% larger than said first number.
- 70. A method in accordance with Claim 58 wherein said method includes the step of calculating the hairpin characteristics of said candidate oligonucleotide sequence.
- 71. A method in accordance with Claim 70 wherein the step of calculating the hairpin characteristics of said candidate oligonucleotide sequence includes the steps of: calculating a complementary sequence to the candidate oligonucleotide sequence by reversing the base pair order of the candidate oligonucleotide sequence and substituting complementary base pairs;

comparing each character of said original candidate oligonucleotide sequence and said complementary sequence;

finding the longest match between said original candidate oligonucleotide sequence and said complementary sequence; and

saving the match with the longest hairpin distance if any two matches have the same length.

72. A method in accordance with Claim 58 wherein said fixed-length target gene subsequences are calculated by a method comprising the steps of:

locating the origin of said subsequence in a set position of said target gene sequence in said preparation file;

cutting a subsequence that is a fixed-length long every preselected number of positions of said target gene sequence in said preparation file; and

sorting said subsequences in said preparation file in lexicographical order beginning at a set position.

73. A method in accordance with Claim 72 wherein the origin of said subsequence is located at position 40 of said target sequence in said preparation file.

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74. A method in accordance with Claim 58 wherein said fixed-length subsequences are calculated by a method comprising the steps of:

locating the origin of said subsequence in the 40th position of said target gene sequence in said preparation file;

cutting a subsequence that is 96 base pairs long of said target gene sequence in said preparation file; and

sorting said subsequences in said preparation file in lexicographical order beginning at a set position.

- 75. A method in accordance with Claim 58 wherein said method includes the step of prepending said preparation file subsequences with identifiers for the sources of each subsequence.
- 76. A method in accordance with Claim 58 wherein said method includes the step of calculating an candidate oligonucleotide sequence's melting temperature comprising:

solving the formula Tm = 81.5 - 16.6(log[Na]) - .63%(formamide) + ((.41 (%(G + C)) - 600)/N);

wherein log[Na] is the sodium concentration, %(G + C) is the fraction of matched base pairs which are G-C complementary, N is the sequence length; and wherein the number of mismatches is equal to zero.

- 77. A method in accordance with Claim 58 wherein said method includes the step for reducing a candidate oligonucleotide sequence's calculated melting temperature by a preselected amount for each percent of mismatch between the candidate oligonucleotide sequence and a user-selected target gene sequence based upon the assumption that there are an equal number of GC and AT base pair mismatches.
- 78. A method in accordance with Claim 58 wherein said method includes the step for reducing a candidate oligonucleotide sequence's calculated melting temperature by a preselected amount comprising the steps of:

reducing said calculated melting temperature by 2 degrees Celsius if an AT mismatch exists; and

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reducing said calculated melting temperature by 4 degrees Celsius if a GC mismatch exists.

79. A method for designing candidate oligonucleotide sequences for use with a gene sequence data source comprising the steps of:

introducing user-selected gene sequence and a user-specified sequence length into a computer system;

storing said gene sequence and said sequence length in the memory of the computer system;

accessing gene sequence data from said gene sequence data source;

accessing the gene sequence source to introduce the user-selected set of gene sequence data and a user-selected set of target gene sequence data from said gene sequence data source into the computer system;

comparing said gene sequences against said target gene sequences employing said criteria;

calculating candidate oligonucleotide sequences of said sequence length that are either common to a pool of user-specified gene sequences or specific to a particular user-specified gene sequence;

calculating the homology between the candidate oligonucleotide sequences and said gene sequence data;

displaying in multiple dimensions the gene sequences which result from the comparisons and calculations characterized in that said display format exhibits:

the starting position of each candidate oligonucleotide sequence in one dimension;

a candidate oligonucleotide sequence's specificity to the target gene sequence in a second dimension; and

superimposed melting temperatures of gene sequences in contrasting presentations in at least an apparent third dimension.

- 80. A method in accordance with Claim 79 wherein said method includes the step of calculating a candidate oligonucleotide sequence's hairpin characteristics.
- 81. A method in accordance with Claim 80 wherein said step of calculating hairpin characteristics for a gene sequence comprises:

calculating a complementary sequence to the said gene sequence by reversing the base pair order of the gene sequence and substituting complementary base pairs;

comparing each character of said original gene sequence and said complementary sequence;

finding the longest match between said original gene sequence and said complementary sequence; and

saving the match with the longest hairpin distance if any two matches have the same length.

82. A method in accordance with Claim 79 wherein the step of displaying further includes producing a cursor moveable along one dimension of said display that selects a position for an expansion of data representing the homology between the candidate oligonucleotide sequences and said gene sequence data; and

displaying in alphanumeric form the homology between the candidate oligonucleotide sequences and said gene sequence data.

- 83. A method in accordance with Claim 79 wherein said display format data is outputted to a printing means.
- 84. A method in accordance with Claim 79 wherein said display format data is saved to a data file.
- 85. A method in accordance with Claim 79 wherein said display format data is exported to another computer system.
- 86. A method in accordance with Claim 79 wherein the step of displaying further includes producing a cursor moveable along one dimension of said display that selects a position for an expansion of data representing the homology between the candidate oligonucleotide sequences and said gene sequence data;

positioning said moveable cursor to select and save particular candidate oligonucleotide sequence information; and

displaying in alphanumeric form the homology between the candidate oligonucleotide sequences and said gene sequence data.

87. A method in accordance with Claim 79 wherein the step of displaying further includes producing a cursor moveable along one dimension of said display that selects a position for an expansion of data representing the homology between the candidate oligonucleotide sequences and said gene sequence data;

positioning said moveable cursor to select and save particular candidate oligonucleotide sequence information;

capturing candidate oligonucleotide sequence information at the user-selected point and storing said information in said memory means; and

displaying in alphanumeric form the homology between the candidate oligonucleotide sequences and said gene sequence data.

88. A method in accordance with Claim 79 wherein said method of displaying comprises:

calculating display output ranges; converting said output ranges to a logarithmic scale; interpolating said converted values; creating a bitmap of said interpolations; and displaying said bitmap on a display device.

89. A method in accordance with Claim 79 wherein said method of displaying comprises:

converting said result values to pixels;

filling a pixel array with said pixels;

performing a binary search into said pixel array;

determining the number of pixels per candidate oligonucleotide sequence to be displayed;

interpolating said pixels at the value of pixels per position minus one; computing an array of said pixel array; and plotting the results on a display device.

90. A method to determine hybridization strength between two or more gene sequences for use with a gene sequence data source, comprising the steps of:

accessing gene sequence data from said gene sequence data source;

comparing base pairs of a first gene sequence and a second gene sequence to determine if a match exists;

incrementing said first gene sequence's bound strength by some first number if a base pair character in said first gene sequence and said second gene sequence match and the matched base pair is equal to a combination of the bases Guanine (G) and Cytosine (C);

incrementing said first gene sequence's bound strength by some second number if a base pair character in said first gene sequence and said second gene sequence match and the matched base pair is equal to a combination of the bases Adenine (A) and Thymine (T);

decrementing said first gene sequence's bound strength by a third number if there is no match in base pairs between said first gene sequence and said second gene sequence;

comparing said first gene sequence's bound strength to said first gene sequence's unbound strength;

setting said first gene sequence's unbound strength equal to its bound strength if said first gene sequence's bound strength is greater than said first gene sequence's unbound strength; and

resetting said first gene sequence's bound strength to zero if said first gene sequence's unbound strength is less than zero.

- 91. A method in accordance with Claim 90 wherein said first and second numbers are greater than zero.
- 92. A method in accordance with Claim 90 wherein said second number is in the order of 42% of said first number.
- 93. A method in accordance with Claim 90 wherein said third number is in the order of 5% larger than said first number.
- 94. A method of calculating the minimum length of any matching gene subsequence comprising:

introducing a user-selected maximum number of mismatches and a user-selected minimum candidate oligonucleotide sequence length;

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subtracting said maximum number of mismatches from said minimum candidate oligonucleotide sequence length to give a first result;

dividing said first result by said maximum number of mismatches plus one to give a second result;

incrementing said second result by one if the remainder is not equal to zero to give a third result; and

truncating said third result to an integer.

95. A method of calculating hairpin characteristics for a gene sequence comprising:

calculating a complementary sequence to the said gene sequence by reversing the base pair order of the gene sequence and substituting complementary base pairs;

comparing each character of said original gene sequence and said complementary sequence;

finding the longest match between said original gene sequence and said complementary sequence; and

saving the match with the longest hairpin distance if any two matches have the same length.

96. A method of creating a preparation file from a user-selected set of target gene sequence data comprising:

cutting said target gene sequence data into fixed-length subsequences; and storing said subsequences in a preparation file.

97. A method of creating a preparation file from a user-selected set of target gene sequence data comprising:

cutting said target gene sequence data into fixed-length subsequences in the order of 96 base pairs in length; and

storing said subsequences in a preparation file.

98. A method in accordance with Claim 97 wherein said fixed-length subsequences are calculated by a method comprising the steps of:

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locating the origin of said subsequence in a set position of said target gene sequence in said preparation file;

cutting a subsequence that is a fixed-length long every preselected number of positions of said target gene sequence in said preparation file; and

sorting said subsequences in said preparation file in lexicographical order beginning at a set position.

99. A method in accordance with Claim 97 wherein said fixed-length subsequences are calculated by a method comprising the steps of:

locating the origin of said subsequence in a set position of said target gene sequence in said preparation file wherein the origin of said subsequence is located at position 40 of said target sequence in said preparation file;

cutting a subsequence that is a fixed-length long every preselected number of positions of said target gene sequence in said preparation file; and

sorting said subsequences in said preparation file in lexicographical order beginning at a set position.

100. A method in accordance with Claim 97 wherein said fixed-length subsequences are calculated by a method comprising the steps of:

locating the origin of said subsequence the 40th position of said target gene sequence in said preparation file;

cutting a subsequence that is 96 base pairs long of said target gene sequence in said preparation file; and

sorting said subsequences in said preparation file in lexicographical order beginning at a set position.

101. A method of forming lists of screens of target gene sequence data comprising:

introducing a user-selected screening threshold; and

forming subsequences of said target gene sequence data of length equal to a user-selected screening threshold.

102. A method of preprocessing a user-selected set of target gene sequence data comprising the steps of:

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searching for sequences without introns in said target gene sequences; extracting target gene sequences that do not contain introns; and storing said extracted target gene sequences.

AMENDED CLAIMS

[received by the International Bureau on 4 April 1994 (04.04.94); original claim 69 amended; remaining claims unchanged (1 page)]

- 68. A method in accordance with Claim 66 wherein said second number is in the order of 42% of said first number.
- 69. A method in accordance with Claim 66 wherein said third number is in the order of 5% larger than said first number.
- 70. A method in accordance with Claim 58 wherein said method includes the step of calculating the hairpin characteristics of said candidate oligonucleotide sequence.
- 71. A method in accordance with Claim 70 wherein the step of calculating the hairpin characteristics of said candidate oligonucleotide sequence includes the steps of:

calculating a complementary sequence to the candidate oligonucleotide sequence by reversing the base pair order of the candidate oligonucleotide sequence and substituting complementary base pairs;

comparing each character of said original candidate oligonucleotide sequence and said complementary sequence;

finding the longest match between said original candidate oligonucleotide sequence and said complementary sequence; and

saving the match with the longest hairpin distance if any two matches have the same length.

72. A method in accordance with Claim 58 wherein said fixed-length target gene subsequences are calculated by a method comprising the steps of:

locating the origin of said subsequence in a set position of said target gene sequence in said preparation file;

cutting a subsequence that is a fixed-length long every preselected number of positions of said target gene sequence in said preparation file; and

sorting said subsequences in said preparation file in lexicographical order beginning at a set position.

73. A method in accordance with Claim 72 wherein the origin of said subsequence is located at position 40 of said target sequence in said preparation file.

FIG. 1



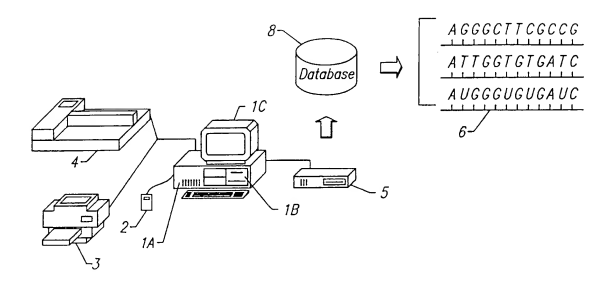
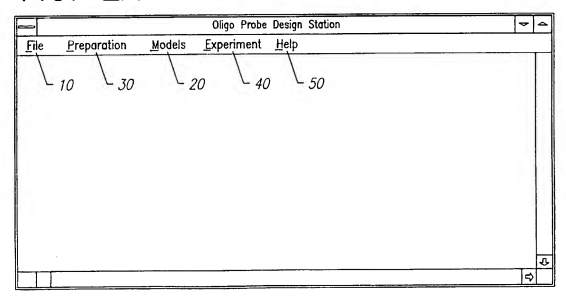
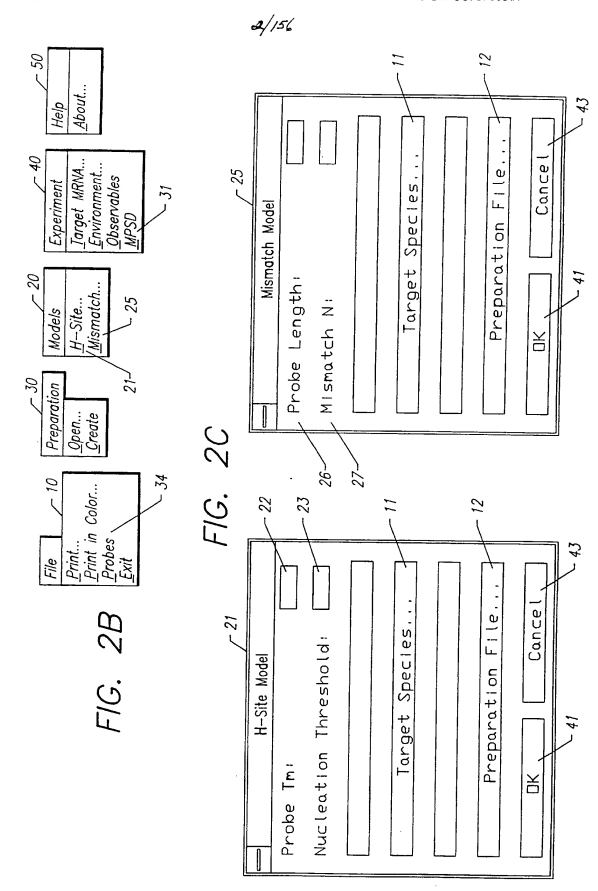
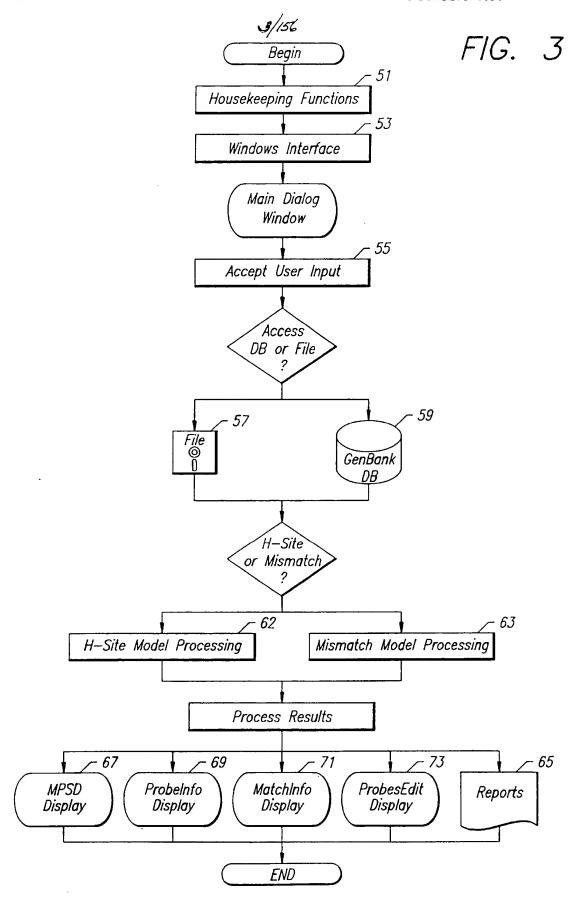


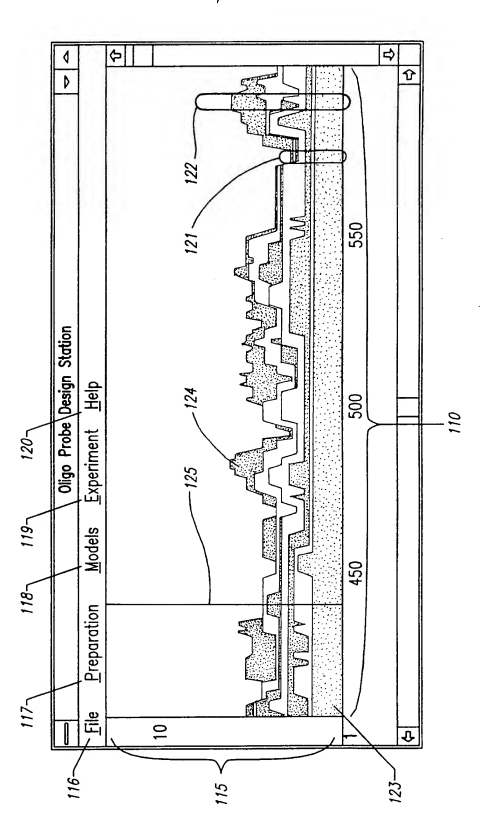
FIG. 2A



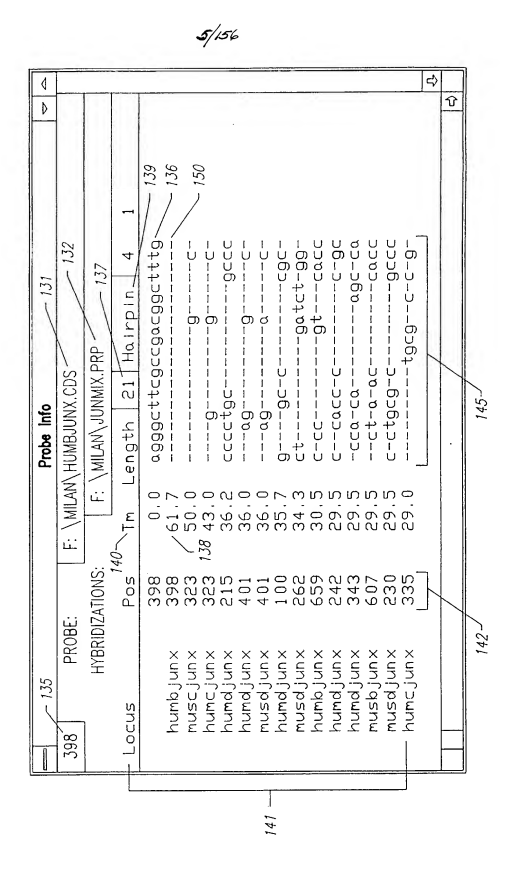




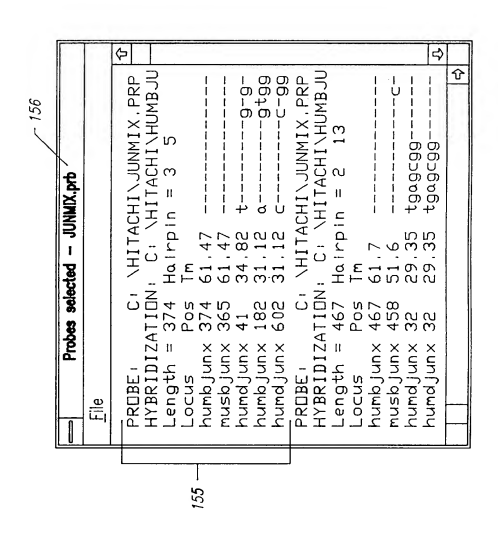
F/G. 4



6.5



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1/15% FIG. 6A (1)

```
PROBE: C:\HITACHI\JUNMIX.PRP
HYBRIDIZATION: C:\HITACHI\HUMBJUNX.CDS
Length = 374 Hairpin = 35
        Pos Tm
Locus
humbjunx 374 61.47 -----
musbjunx 365 61.47 -----
            34.82 t-----g-g--agt
humdjunx 41
humbjunx 182 31.12 a----gtgg--gc
humdjunx 602 31.12 c----c-ggg-gc
humdjunx 602 31.12 c----c-ggg-gc
PROBE: C:\HITACHI\JUNMIX.PRP
HYBRIDIZATION: C:\HITACHI\HUMBJUNX.CDS
Length = 377 Hairpin = 214
Locus
        Pos Tm
humbjunx 377 61.55 -----
musbjunx 368 61.55 -----
humdjunx 383 28.12 tg-cg-c--g-----
musdjunx 383 28.12 tg-ca-c--g-----
musdjunx 383 28.12 tg-ca-c--q-----
PROBE: C:\HITACHI\JUNMIX.PRP
HYBRIDIZATION: C:\HITACHI\HUMBJUNX.CDS
Length = 389 Hairpin = 33
Locus
        Pos Tm
humbjunx 389 61.7
muscjunx 314 56.65 -c-----
musbjunx 380 50.85 -----t--q
humcjunx 314 49.35 -t----g-----
humdjunx 395 33.85 -----tt-qc--aq
musdjunx 395 33.85 -----tt-qc--aa
humcjunx 326 32.35 q-ttcgcc----tq
humdjunx 404 32.35 -- ttcgcc-----t-
muscjunx 326 32.35 gcttcgcc----tg
musdjunx 253 30.85 gacg-gct-ct-----
humbjunx 953 30.65 g----t-c-cagct-
musdjunx 83 27.3 cc-gcggt-gt-----g
```

FIG. 6A (2)

```
PROBE: C:\HITACHI\JUNMIX.PRP
HYBRIDIZATION: C:\HITACHI\HUMBJUNX.CDS
Length = 397 Hairpin = 41
Locus
        Pos Tm
humbjunx 397 61.55 ----
muscjunx 322 53.44 -----q---
humcjunx 322 45.33 ----g----g---
musbjunx 388 41.38 -----t--g----t
humdjunx 214 36.83 cccctgc-----
humdjunx 99
            36.16 cg----gc-c-----
musdjunx 261 34.55 -ct-----gatct
humdjunx 400 33.27 c---ag-----g---
musdjunx 400 33.27 c---ag----a---
humcjunx 334 32.28 -----tgcg--c-
humdjunx 412 32.28 -----t-a-q-c-
muscjunx 334 32.28 -----tgcq--c-
humbjunx 658 30.17 cc-cc----gt---
humdjunx 241 28.95 -c--cacc-c----
humdjunx 342 28.95 c-cca-ca----aq
musbjunx 606 28.95 ---ct-a-ac-----
musdjunx 229 28.95 -c-ctqcq-c----
musdjunx 91
            26.67 -gt-----gcc-ccg
PROBE: C:\HITACHI\JUNMIX.PRP
HYBRIDIZATION: C:\HITACHI\HUMBJUNX.CDS
Length = 417 Hairpin = 215
        Pos Tm
Locus
humbjunx 417 60.08 ---
musbjunx 408 55.52 ----
humdjunx 420 37.3 c----g---t-a-
musbjunx 61
            29.0
                 g---gg-----ca-cctgt-
muscjunx 672 26.27 gc-gc----a-g--aga--
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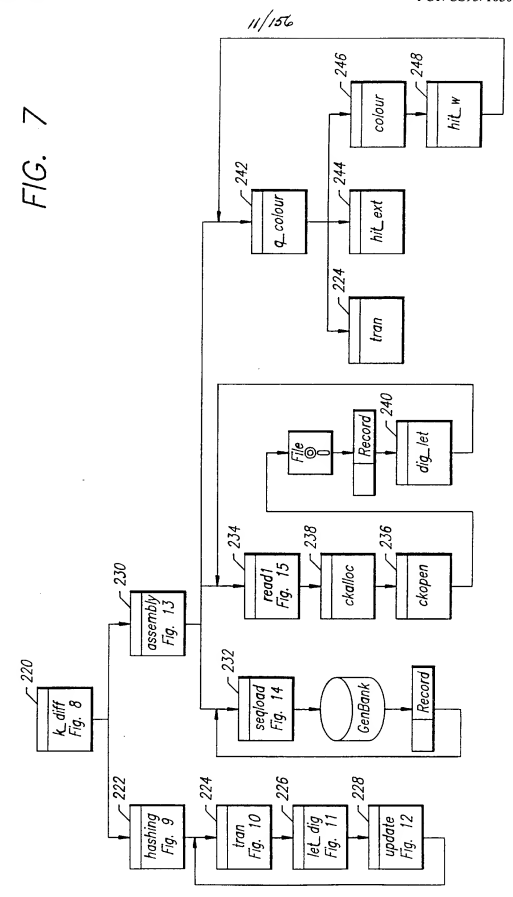
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9/156 FIG. 6A (3)

			JUNMIX.PRP
			HITACHI\HUMBJUNX.CDS
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Locus	Pos	Tm	
humbjunx	461	61.63	
musbjunx	452	61.63	
musbjunx	452	61.63	
PROBE: C	:\HI'	TACHI\	JUNMIX.PRP
HYBRIDIZ	ATIO	N: C:\]	HITACHI\HUMBJUNX.CDS
Length =	467	Hairp.	in = 2 13
Locus	Pos	Tm	
humbjunx	467	61.7	
musbjunx	458	51.6	c-g
humdjunx	32	29.35	tgagcgggcgg
humdjunx	32	29.35	tgagcgggcgg
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HYBRIDIZA	OITA	V: C:/I	HITACHI\HUMBJUNX.CDS
Length =			in = 2 4
Locus	Pos	Tm	
humbjunx	477	61.37	
humdjunx	489	34.93	c-ccg
humdjunx	489	34.93	c-ccg
			JUNMIX.PRP
			HITACHI\HUMBJUNX.CDS
Length =			in = 3 3
Locus	Pos	$\mathbf{T}\mathfrak{m}$	
humbjunx	487	61.14	
musdjunx	74	51.0	ct
humdjunx	499	45.64	tg
humdjunx	527	30.72	CC-C-C
musdjunx	97	30.72	ttc-cg
musdjunx	580	30.72	-cct-g
musdjunx	637	30.72	cc-ccg
musdjunx	637	30.72	cc-ccg

FIG. 6A (4)

PROBE: C:\HITACHI\JUNMIX.PRP
HYBRIDIZATION: C:\HITACHI\HUMBJUNX.CDS
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Locus Pos Tm
humbjunx 498 61.26 -----humbjunx 498 61.26 -----PROBE: C:\HITACHI\JUNMIX.PRP
HYBRIDIZATION: C:\HITACHI\HUMBJUNX.CDS
Length = 504 Hairpin = 3 2
Locus Pos Tm
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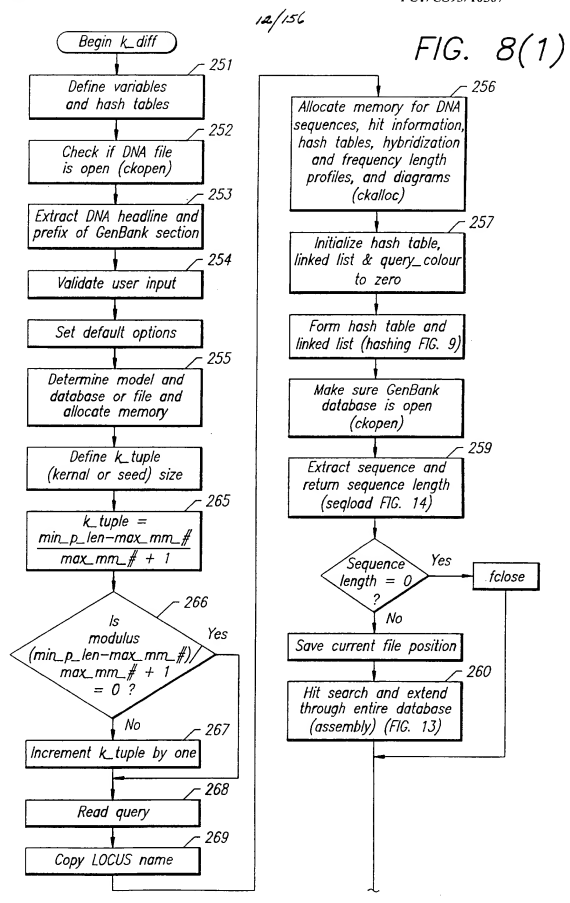
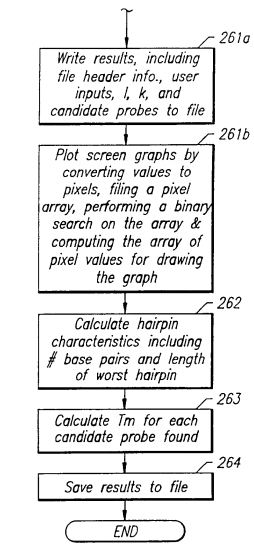
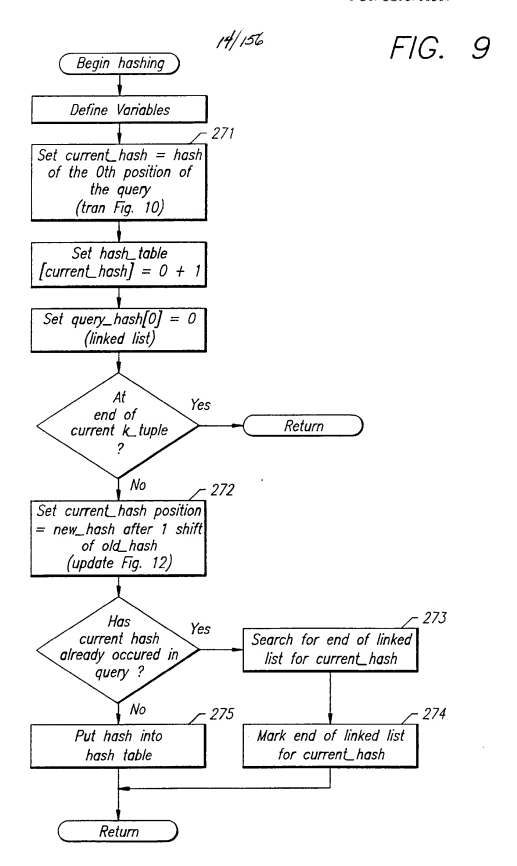


FIG. 8(2)





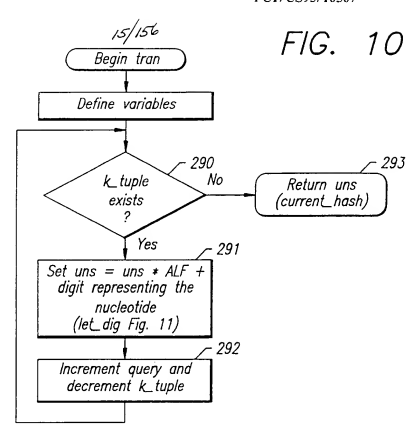
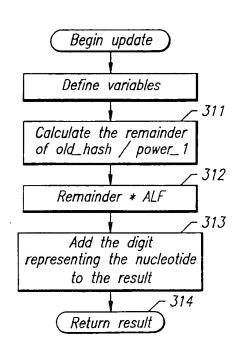
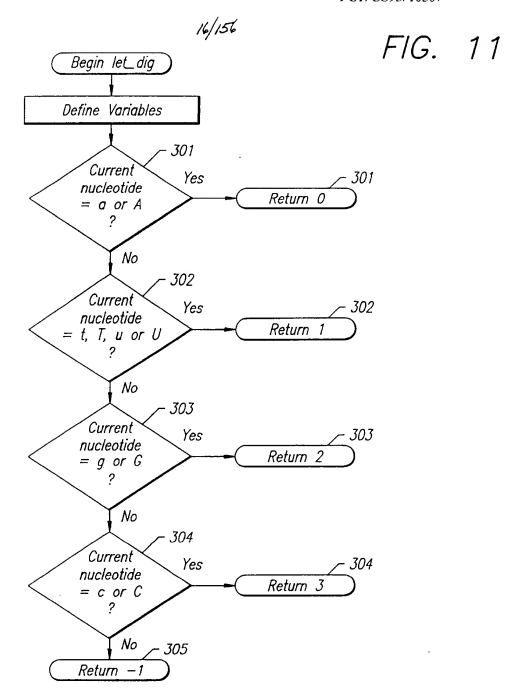
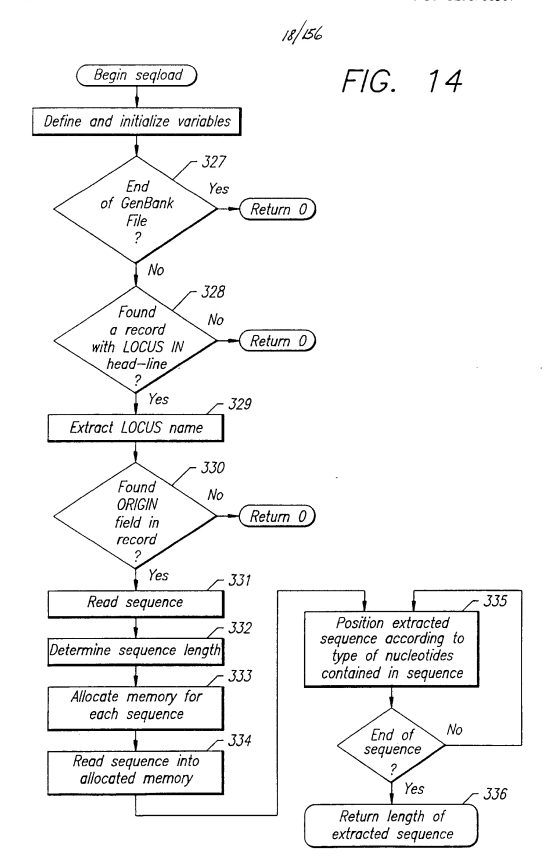


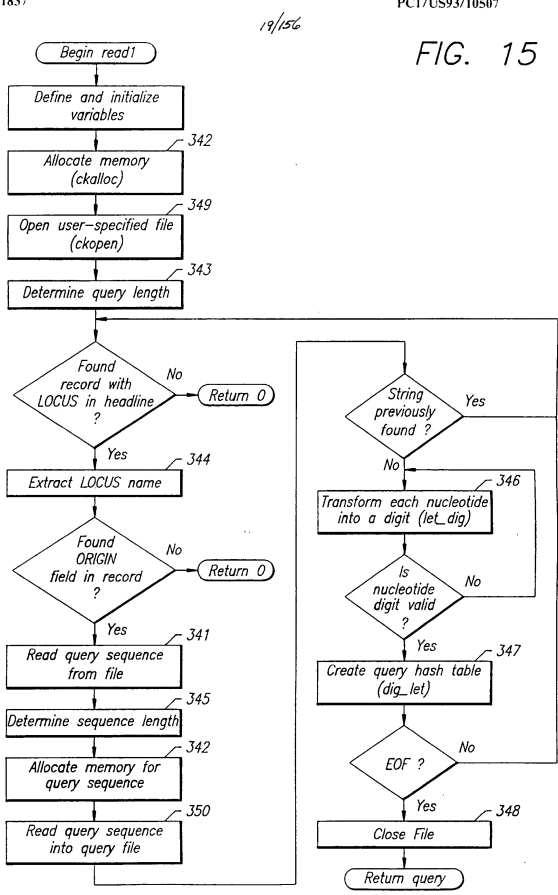
FIG. 12



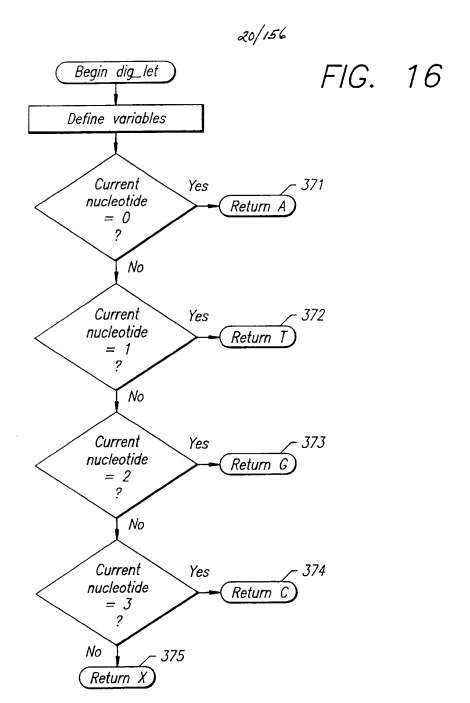


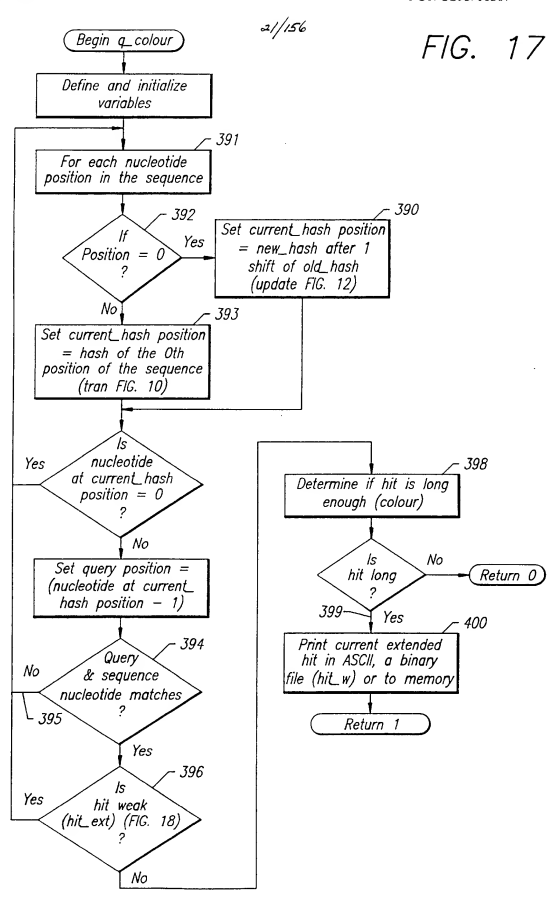
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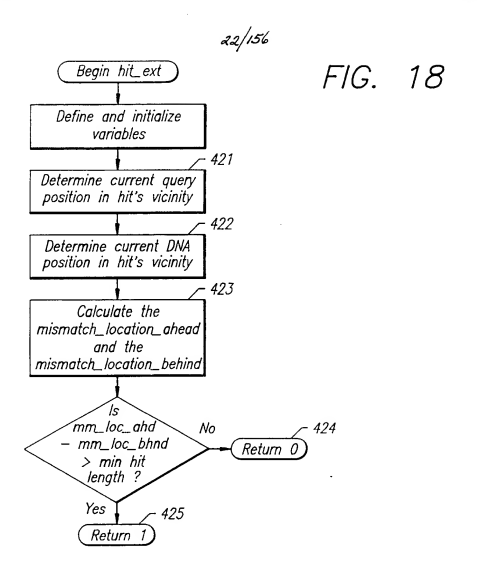


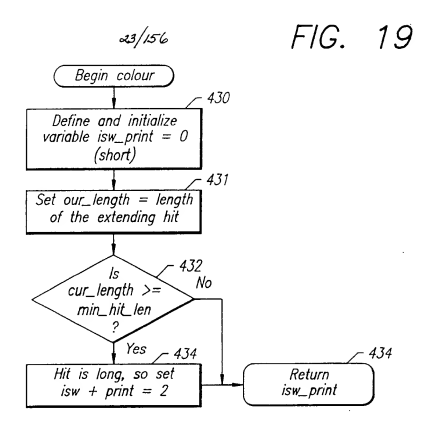
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ω	DesignStation	<pre>C:\HITACHI\HUMBJUNX.CDS : C:\HITACHI\JUNMIX.SEQ</pre>
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_		႕	٦	Н	႕	TGCA	TAAAA	TGGA	TGCACTAAAATGGAACAGCCC	
• •		႕	Н	Н	Н	GCACT	GCACTAAAATGGAACAGCC	GGAA	CAGCCCT	
		⊣	Н	Н	Н	CACTA	AAAATG	GAACI	CACTAAAATGGAACAGCCCTT	
•	_	⊣	Н	Н	Н	ACTA	AAATGG	:AACA	ACTAAAATGGAACAGCCCTTC	
ν-,		٦	۲	Н	Н	CTAA	AATGGA	ACAG	CTAAAATGGAACAGCCCTTCT	
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• •	7	۲	႕	Н	Н	AAATC	SGAACA	GCCC.	AAATGGAACAGCCCTTCTACC	
• •	_	۲	٦	႕	ᠳ	AATGO	SAACAG	CCCT	AATGGAACAGCCCTTCTACCA	
	1	⊣	٦	႕	Н	ATGG1	AACAGC	CCTT	ATGGAACAGCCCTTCTACCAC	
•		۲.	۲	٦	႕	TGGA	TGGAACAGCCCTTCTACCA	CTTC	FACCACG	

FIG. 20 (2)

GGAACAGCCCTTCTACCACGA	GAACAGCCCTTCTACCACGAC	AGCCCTTCTACCACGA	AGCCCTTCTACCACGAC	TACCACGACG	PACCACGACGAC	GCCCTTCTACCACGACGACTC	\mathbf{c}	CTCA	CTTCTACCACGACGACTCATA	TACCACGACGACTCAT	ATAC	TACA	TACCACGACGACTCATACACA	AC	Ø	ACAGC	CACAGC	ACAGCT	CAGCTAC	AGCTACG	CAGCTACGG
Н	Н	٦	۲	СЧ	٦	٦	Н	Н	Н	Н	Н	Н	Н	Н	Н	~	щ	Н	٦	Н	۲
⊣	Н	Н	Н	Н	ᠳ	7	Н	Н	7	٦	Н	Н	7	ਜ	٦	ᆫ	Н	1	ᠬ	Н	1
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GACTCATACACAGCTACGGGA	ACTCATACACAGCTACGGGAT	CTCATACACAGCTACGGGATA	TCATACACAGCTACGGGATAC	CATACACAGCTACGGGATACG	ATACACAGCTACGGGATACGG	TACACAGCTACGGGATACGGC	ACACAGCTACGGGATACGGCC	CACAGCTACGGGATACGGCCG	ACAGCTACGGGATACGGCCGG	CAGCTACGGGATACGGCCGGG	AGCTACGGGATACGGCCGGGC	GCTACGGGATACGGCCGGGCC	CTACGGGATACGGCCGGGCCC	TACGGGATACGGCCGGGCCCC	ACGGGATACGGCCCGGGCCCCT	CGGGATACGGCCGGGCCCCTG
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FIG. 20 (4)

GGGATACGGCCCGGGCCCCTGG	Ü	F⊢	ATACGGCCGGGCCCCTGGTGG	TACGGCCGGCCCCTGGTGGC	ACGGCCGGGCCCCTGGTGGCC	CGGCCGGCCCCTGGTGGCCT	CCCCCCCTCCTCCTC	GCCGGGCCCTGGTGGCCTCT	CCGGGCCCCTGGTGGCCTCTC	GCCTCT	GGGCCCCTGGTGGCCTCTCTC	GGCCCCTGGTGGCCTCTCTCT	Σ	CCCCTGGTGGCCTCTCTAC	CCCTGGTGGCCTCTCTACA	CCTGGTGGCCTCTCTCTACAC	TCTACAC	TCTACAC	TACACGA	CACGAC
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TGGCCTCTCTACACGACTA	GGCCTCTCTACACGACTAC	GCCTCTCTACACGACTACA	CCTCTCTACACGACTACAA	CTCTCTCTACACGACTACAAA	TCTCTCTACACGACTACAAAC	CTCTCTACACGACTACAAACT	TCTCTACACGACTACAAACTC	CTCTACACGACTACAAACTCC	TCTACACGACTACAAACTCCT	CTACACGACTACAAACTCCTG	TACACGACTACAAACTCCTGA	ACACGACTACAAACTCCTGAA	CACGACTACAAACTCCTGAAA	ACGACTACAAACTCCTGAAAC	CGACTACAAACTCCTGAAACC	GACTACAAACTCCTGAAACCG	ACTACAAACTCCTGAAACCGA	CTACAAACTCCTGAAACCGAG	TACAAACTCCTGAAACCGAGC	ACAAACTCCTGAAACCGAGCC
Н	Н	H	Н	Н	Н	Н	Н	۲	٦	Н	Н	Н	Н	7	Н	٦	Н	۲	۲	٦
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CAAACTCCTGAAACCGAGCCT	AAACTCCTGAAACCGAGCCTG	AACTCCTGAAACCGAGCCTGG	ACTCCTGAAACCGAGCCTGGC	CTCCTGAAACCGAGCCTGGCG	TCCTGAAACCGAGCCTGGCGG	CCTGAAACCGAGCCTGGCGGT	CTGAAACCGAGCCTGGCGGTC	TGAAACCGAGCCTGGCGGTCA	GAAACCGAGCCTGGCGGTCAA	AAACCGAGCCTGGCGGTCAAC	CTGGC	ACCGAGCCTGGCGGTCAACCT	CCGAGCCTGGCGGTCAACCTG	CGAGCCTGGCGGTCAACCTGG	GAGCCTGGCGGTCAACCTGGC	AGCCTGGCGGTCAACCTGGCC	GCCTGGCGGTCAACCTGGCC	CCTGGCGGTCAACCTGGCCGA	SGCGGTCA A CTGGCGG	<u></u>	
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Н	Н	Н	Н	Н	Н	⊣	Н	-	Н	٦	٦	7	7	Н	٦	٦	ᡤ	Н	٦	н	
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FIG. 20 (7)

GGCGGTCAACCTGGCCGACCC	GCGGTCAACCTGGCCGACCCC	CGGTCAACCTGGCCGACCCCT	GGTCAACCTGGCCGACCCCTA	GTCAACCTGGCCGACCCCTAC	TCAACCTGGCCGACCCCTACC	CAACCTGGCCGACCCCTACCG	AACCTGGCCGACCCCTACCGG	ACCTGGCCGACCCCTACCGGA	CCTGGCCGACCCCTACCGGAG	CTGGCCGACCCCTACCGGAGT	TGGCCGACCCTACCGGAGTC	GGCCGACCCTACCGGAGTCT	GCCGACCCTACCGGAGTCTC	CCGACCCCTACCGGAGTCTCA	CGACCCCTACCGGAGTCTCAA	GACCCCTACCGGAGTCTCAAA	ACCCCTACCGGAGTCTCAAAG	CCCCTACCGGAGTCTCAAAGC	CCCTACCGGAGTCTCAAAGCG	CCTACCGGAGTCTCAAAGCGC
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CGK00035725

CTACCGGAGTCTCAAAGCGCC	TACCGGAGTCTCAAAGCGCCT	ACCGGAGTCTCAAAGCGCCTG	CCGGAGTCTCAAAGCGCCTGG	CGGAGTCTCAAAGCGCCTGGG	GGAGTCTCAAAGCGCCTGGGG	GAGTCTCAAAGCGCCTGGGGC	AGTCTCAAAGCGCCTGGGGCT	GTCTCAAAGCGCCTGGGGCTC	TCTCAAAGCGCCTGGGGCTCG	CTCAAAGCGCCTGGGGCTCGC	TCAAAGCGCCTGGGGCTCGCG	CAAAGCGCCTGGGGCTCGCGG	AAAGCGCCTGGGGCTCGCGGA	AAGCGCCTGGGGCTCGCGGAC	AGCGCCTGGGGCTCGCGGACC	GCGCCTGGGGCTCGCGGACCC	\mathcal{O}	GCCTGGGGCTCGCGGACCCGG	CCTGGGGCTCGCGGACCCGGC	CTGGGGCTCGCGGACCCGGCC
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Н	Н	Ч	7	Ч	Н	J	Н	러	٢	Н	Ч	٦	Н	Н	٦	7	7	٢	Н	-
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FIG. 20 (9)	1 GGGCTCGGGACCCGGCCCA 1 GGGCTCGCGACCCGGCCCAGA 1 GGCTCGCGGACCCGGCCCAGA 1 GCTCGCGGACCCGGCCCAGAG 1 CTCGCGGACCCGGCCCAGAGG 1 TCGCGGACCCGGCCCAGAGGG 1 CCGGACCCGGCCCAGAGGGC 1 GCGGACCCGGCCCAGAGGGC 1 GCGGACCCGGCCCAGAGGGCG 1 GCGGACCCGGCCCAGAGGGCG 1 GCGGACCCGGCCCAGAGGGCG 1 CCGGCCCAGAGGGCGGCG 1 CCGGCCCAGAGGGCGGCG 1 CCGGCCCAGAGGGCGGCG 1 CCCGGCCCAGAGGGCGGCGGCG 1 CCCGGCCCAGAGGGCGGCGGCGCGCGCGCGCGCGCGCGC
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	4.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4
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FIG. 20 (10)

GAGGCGGCGGTGGCGGCAGC	Ū	GGGCGGCGGTGGCGGCAGCTA	TGGCGGCAGC	゚゙	SGCAGCTAC	LACT	GCGGTGGCGGCAGCTACTTTT	CGGTGGCGGCAGCTACTTTTC	CTTTTC	CTACTTTC	CTTTTCTG	G	CTGGT	CGGCAGCTACTTTTCTGGTCA	Ø	GCAGCTACTTTCTGGTCAGG	CTGGTCAG	CTGGTCAGGG	TGGTCAGGGC	TCTGGTCAGGGCT
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J	7	Ч	Н	Н	Ч	٦	IJ	႕	٦	Н	Н	Н	Н	႕	, - 	Н	Н	႕	Н	Н
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81	82	183	184			187	88	89						95		-97	98	66	00	01

FIG. 20 (11)

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	CTGGTCAGGGCTCG	TGGTCAGGGCTCGG	GGTCAGGGCTCGGA	GTCAGGGCTCGGAC	TCAGGGC	TCAGGGCTCGGACAC	AGGGCTCGGACACC	TCAGGGCTCGGACACCG	GGCTCGGACACCGG	TCAGGGCTCGGACACCGGC	TCAGGGCTCGGACACCGGCG	AGGGCTCGGACACCGGCGC	GGGCTCGGACACCGGCGCG	AGGGCTCGGACACCGGCGCGT	GCTCGGACACCGGCGCGTC	CACCGGCGCGT	CACCGGCGCGTCTC	CCGGCG	GCGCGTCTCT	CCGGCGCGTCTCTCA	GGCGCGTCTCTCAA
•	TACTTTT	ACTTTTC	CTTTTCT	TTTTCTG	TTTCTGG	TTCTGGT	TCTGGTC	CTGGTCA	TGGTCAGGG	GGTCAGG	GTCAGGG(TCAGGGC	CAGGGCT	AGGGCTC	GGGCTCG	GGCTCGGA	GCTCGGA	CTCGGACA	TCGGACACCG	CGGACAC	GGACACC
	Н	⊣	Н	Н	Н	Н	Н	Ч	Н	Н	Н	Н	~	Н	بر	Н	٦	Н	⊣	-	Н
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FIG. 20 (12)

GACACCGGCGCGTCTCTCAAG	ACACCGGCGCGTCTCTCAAGC	CACCGGCGCGTCTCTCAAGCT	ACCGCCGCGTCTCTCAAGCTC	CCGCCCCTCTCAAGCTCG	CGGCGCGTCTCTCAAGCTCGC	GGCGCGTCTCTCAAGCTCGCC	GCGCGTCTCTCAAGCTCGCCT	CGCGTCTCTCAAGCTCGCCTC	GCGTCTCAAGCTCGCCTCT	CGTCTCAAGCTCGCCTCTT	GICTCTCAAGCTCGCCTCTTC	TCTCTCAAGCTCGCCTCTTCG	CTCTCAAGCTCGCCTCTTCGG	TCTCAAGCTCGCCTCTTCGGA	CTCAAGCTCGCCTCTTCGGAG	TCAAGCTCGCCTCTTCGGAGC	CAAGCTCGCCTCTTCGGAGCT	AAGCTCGCCTCTTCGGAGCTG	AGCTCGCCTCTTCGGAGCTGG	GCTCGCCTCTTCGGAGCTGGA	CTCGCCTCTTCGGAGCTGGAA
Н	Н	Н	. , –	٦	Н	Н	Н	٦	Н	٦	٦	۲	Н	Ч	Н	Н	٦	7	7	٦	٦
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7	Н	Н	Т	П	Н	른	Н	Н	٦	⊣	٦	Н	⊣	٦	1	7	Т	Н	٦	ᠬ	Н
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	224			227					$^{\circ}$			\mathfrak{C}	\mathcal{C}			239			242	243	244

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TCGCCTCTTCGGAGCTGGAAC	CTCTTCGGAGCTGGAACG	TCTTCGGAGCTGGAACGC	TGGAACGC			CTGGAACGCCT	TGA	GA	Ę	TGATTG	CTGATTG	TGATTGTC	GATTGTCC	ATTGTCC	CCTGATTGTCCC	CTGATTGTCCCC	TGATTGTCCCCA	SATTGTCCCCAAC	ATTGTCCCCAAC	
TCGCCTC	CGCCTCT	GCCTCTT	CCTCTTC	CTCTTCG	TCTTCGG	CTTCGGAG	TTCGGAG	TCGGAGC	CGGAGCT	GGAGCTG	GAGCTGGAACGC	AGCTGGA	GCTGGAACGCCT	CTGGAACGCCTG	TGGAACG	GGAACGC	GAACGCC	AACGCCT	ACGCCTG	くり上してして
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4	246	4	248	4	Ŋ	251	S	253	വ	S	256	Ŋ	S	Ŋ	9	9	262	263	264	265

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GCCTGATTGTCCCCAACAGCA	CCTGATTGTCCCCAACAGCAA	CTGATTGTCCCCAACAGCAAC	TGATTGTCCCCAACAGCAACG	GATTGTCCCCAACAGCAACGG	ATTGTCCCCAACAGCAACGGC	TTGTCCCCAACAGCAACGGCG	TGTCCCCAACAGCAACGGCGT	GTCCCCAACAGCAACGGCGTG	TCCCCAACAGCAACGGCGTGA	CCCCAACAGCAACGGCGTGAT	CCCAACAGCAACGGCGTGATC	CCAACAGCAACGGCGTGATCA	CAACAGCAACGGCGTGATCAC	AACAGCAACGGCGTGATCACG	ACAGCAACGGCGTGATCACGA	CAGCAACGGCGTGATCACGAC	AGCAACGGCGTGATCACGACG	GCAACGGCGTGATCACGACGA	CAACGGCGTGATCACGACGAC	AACGCCTGATCACGACGACG
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99		89	69	70	71	72	73	74	75	9/	11	78	19	80	81	82	83	84	85	86

FIG. 20 (15)

		IJ	႕	⊣	٦	٦	ACGCCGTGATCACGACGACGC
		~	~	⊣	, H	Н	CGGCGTGATCACGACGACGCC
ω		~	٦	Н	⊢	-	GGCGTGATCACGACGACGCCT
9		⊣	Н	Н	7	٦	GCGTGATCACGACGACGCCTA
		Н	J	Н	ᅥ	٦	CGTGATCACGACGCCTAC
9		\vdash	Ч	\leftarrow	٦	Н	O
293	21	Н	۲	ಗ	٦	٦	TGATCACGACGACGCCTACAC
σ		⊣	Ч	႕	\vdash	٦	AC
		7	, - 	⊣		۲-	ATCACGACGACGCCTACACCC
σ		٦	~	⊣	٦	٦	TCACGACGCCTACACCCC
$\boldsymbol{\omega}$		႕	~	⊣		٦	CACGACGCCTACACCCCC
σ		↤	근	IJ	H	٦	ACGACGCCTACACCCCCG
$\boldsymbol{\sigma}$		٦	┍┥	-	· H	Н	CGACGACGCCTACACCCCCGG
		٦	Н	٦	⊣	Н	GACGACGCCTACACCCCCGGG
0		-	ᡤ	⊣	-	Н	ACGACGCCTACACCCCCGGGA
0		IJ	⊣	~	 1	Н	CGACGCCTACACCCCCGGGAC
_		۲,	⊣	IJ	H	٦	GACGCCTACACCCCCGGGACA
304		۲	7	7	, H	Ч	ACGCCTACACCCCCGGGACAG
302		Н	⊣	7		Н	CGCCTACACCCCCGGGACAGT
306		Η.	⊢	7		Н	GCCTACACCCCCGGGACAGTA
307	21		Ļ	٦	٦	щ	CCTACACCCCCGGGACAGTAC
308	21	Н	, 1	٦	۲	Н	CTACACCCCGGGGACAGTACT

FIG. 20 (16)

TACACCCCCGGGACAGTACTT	ACCCCGGGACAGTAC	ACCCCCGGGACAGTAC	TAC	GTACTTTTA	CCCGGGACAGTACTTTAC	CGGGACAGTACTTTAC	TTTACCC	THTTACC	ACTTTTACC	ACTTTTAC	AGTACTTTACCCC	AGTACTTTACCCCCC	ACTTTACCCCCGC	PTTTACCC	PTTACCCCGC		CTTTTACCCCCC			
Н	-	٦	Н	٦	٦	Н	Н	٦	٦	Н	Н			Н	Ч	Н	T	۲	-	٦
Н	J	ᆸ	Н	H	Н	ત	٦	Н	Н	Н	H	Н	႕	Н	7	Н	Н		Н	Н
Н	٦	٦	⊣	Н	М	Н	٦	۲	٦	ᠳ	٦	Н	Ч	٦	Н	Н	Н	Н	۲	H
Ч	Н	Н	Н	Н	Н	႕	Н	-	J	Н	Н	7	7	را ٔ	7	٦	۲	7	⊣	Н
٦	Н	Н	Н	Н	႕	IJ	Н	⊣	႕	Н	٦	.H	7	٦	7	⊣	٦	Н	Н	7
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
309	\vdash	\leftarrow	312	\vdash	314	315	\vdash	\vdash	٦	\vdash	2	\sim	322	2	\sim	325	7	~	328	329

FIG. 20 (17)

TTACCCCCGGGGGGTGGCAG	C		CCCCCCCGGGGGTGGCAGCGG	CCCCGCGGGGTGGCAGCGGT	CCCGCGGGGTGGCAGCGGTG	CGGT	CGCGGGGGTGGCAGCGGTGGA	GCGGGGGTGGCAGCGGTGGAG	GCGGTGGAG	GGGGGTGGCAGCGGTGGAGGT	$_{ m GT}$	GGGTGGCAGCGGTGGAGGTGC	GGTGGCAGCGGTGGAGGTGCA	GTGGCAGCGGTGGAGGTGCAG	TGGCAGCGGTGGAGGTGCAGG	GGCAGCGGTGGAGGTGCAGGG	A G	GGTGC	GTGC	AGGGGGC
Н	Н	Н	~	~	Н	Н	Н	Н	-	Н	٦	٦	٦	Н		~	7		7	Н
٦	Н	~	-	~	ᆏ	П	Н	Н		러	 1	႕	۲	Н	⊣	~	٦	Н	٦	Н
Н		٦	Н	Н	ᅥ	Н	Н	Н	႕	Н	٦	Ч	Н	႕	Ч	Н	Н	~	٦	႕
Н	~	ㄷ	႕	Н	Н	Н	Н	Н	IJ	Н	Н	⊣	႕	Н	ᠬ	Н	۲	H	7	٦
⊣	1	7	٦	Н	Н	Н	٦	႕	٦	٦	Н	Н	IJ	Н	\vdash	~	Н	⊣	Н	Н
21	. 21	2	~	7	2	7	7	7	7	~	7	7	7	7	2	~	2	~	~	21
330	331	332	333							4	4	4		4		4	347	348	349	350

FIG. 20 (18)

CGGTGGAGGTGCAGGGGGCGC	GGTGGAGGTGCAGGGGGCGCA	GTGGAGGTGCAGGGGGCGCAG	TGGAGGTGCAGGGGGCGCAGG	GGAGGTGCAGGGGGCGCAGGG	GAGGTGCAGGGGGCGCAGGGG	AGGTGCAGGGGGCGCAGGGGG	GGTGCAGGGGGCGCAGGGGGC	GTGCAGGGGGCGCAGGGGGCG	TGCAGGGGGCGCAGGGGGCGG	GCAGGGGGGCAGGGGGGGG	CAGGGGGGCAGGGGGGGGCG	AGGGGCGCAGGGGGCGGCGT	GGGGCGCAGGGGGCGGCGTC	GGGCGCAGGGGCGGCGTCA	GGGCGCAGGGGGCGCGTCAC	GGCGCAGGGGGCGCGTCACC	GCGCAGGGGGCGCGTCACCG	CGCAGGGGGGGGCGTCACCGA	GCAGGGGCGGCGTCACCGAG	CAGGGGGGGGGTCACCGAGG	AGGGGCGCGTCACCGAGGA
٦	_	٢	Н	(Н	٢	4	۲	Ч	Н	٢	Ч	Н	Н	Н	⊣	~	Н	2	2	2
٦	П	۲	ı	٦	Н	⊣	ᆏ	Н	႕	╓┤	Н	Н	H	Н	ч	Н	⊣	Н	2	2	2
Н	Н	Ч	Н	٦	\vdash	٦	П	٦	Н	٦	٦	٦	٦	Н	~	٦	Н	_	7	7	7
Н	러	гH	٦	٦	٦	7	-	J	1	1	1	1	J	1	Н	٦	H	7	2	7	7
٦	٦	٦	٦	٦	Н	٦	٦	Н	~	Н	۲	٦	٦	۲	Н	Н	٦	IJ	Ч	7	7
																4					
21		21					21									21		21		21	21
351	352	353	354	355	356	357	358				362		Ó	365	366	367	368		370	371	372

FIG. 20 (19)

GGGGCGCGTCACCGAGGAG	GGGGCGCGTCACCGAGGAGC	GGGCGCCTCACCGAGGAGCA	GGCGGCGTCACCGAGGAGCAG	GCGGCGTCACCGAGGAGCAGG	CGGCGTCACCGAGGAGCAGGA	GGCGTCACCGAGGAGCAGGAG	GCGTCACCGAGGAGCAGGAGG	CGTCACCGAGGAGCAGGAGGG	GTCACCGAGGAGCAGGGC	TCACCGAGGAGCAGGAGGGCT	CACCGAGGAGCAGGGGCTT	ACCGAGGAGCAGGAGGGCTTC	CCGAGGAGCAGGAGGGCTTCG	CGAGGAGCAGGAGGGCTTCGC	GAGGAGCAGGAGGGCTTCGCC	AGGAGCAGGAGGGCTTCGCCG	GGAGCAGGAGGCTTCGCCGA	GAGCAGGAGGCTTCGCCGAC	AGCAGGAGGCTTCGCCGACG	GCAGGAGGGCTTCGCCGACGG
~	~	2	~	2	2	2	2	~	2	2	2	ന	സ	က	2	2	7	2	2	2
2	7	2	2	2	7	2	2	7	2	2	2	က	က	8	2	2	2	2	2	2
7	~	7	7	~	7	7	7	7	7	7	7	က	സ	က	7	2	2	7	2	7
7	7	2	7	7	8	2	7	2	7	7	~	ო	М	М	2	7	2	7	2	2
2	2	2	2	2	7	2	7	2	2	7	2	⊣	٦	-	-	-	Н	Н	⊣	Н
21	21		21						21		21		21	21			21	21	21	21
373	374		376	•	_	379	380	381	382	383	384	-	386	387	388	389	390	391	392	393

FIG. 20 (20)

	TTCGCCGACGGCT	AGGGCTTCGCCGACGGCTT	CTTCGCCGACGGCTTT	CTT	ACGGCTTTGT	CTTCGCCGACGGCTTTGTC	GCTTCGCCGACGGCTTTGTCA	CGGCTTTGTCAA	CTTTGTCAAA	TCGCCGACGCCTTTGTCAAAG	CGCCGACGCCTTTGTCAAAGC	TGTCAAAGCC	CCGACGCTTTGTCAAAGCCC	TCAAAGCCCT	TCAAAGCCCTG	TGTCAAAGCCCTGG	CAAAGCCCTGGA	TCAAAGCCCTGGAC	AAAGCCCTGGACG	CCCTGGACGA
CAGGAGGGCTTCGCCGACGG	AGGAGGGCTT	GGAGGCCTTC	GAGGGCTTCG	AGGGCTTCGCCGACGG	GGGCTTCGCCGACGGCTTT	GGCTTCGCCG	GCTTCGCCGA	CTTCGCCGAC	TTCGCCGACGG	TCGCCGACGG	CGCCGACGGC	GCCGACGGCTT	CCGACGGCTT	CGACGCCTTTG	GACGGCTTTG	ACGCCTTTGT	CGGCTTTGTC	GGCTTTGTCA	GCTTTGTCAA	CTTTGTCAAAGCC
7	2	7	_		٦	⊣	~	Н	⊣	٦	Н	Н	Н	⊣	7	7	7	7	2	7
7	7	7	П	7	Ч	۲	Н	႕	႕	Ч	٦	Н	۲	⊣	7	2	2	2	7	2
2	2	7	⊣	٦	႕	႕	႕`	Т	7	٦	٦	7	٦	7	7	7	7	2	7	7
2	2	7	႕	٦	٦	H	7	႕	۲	႕	႕	ᆏ	႕	Н	7	7	2	7	2	7
	႕	Н	Н	႕	⊣	٦	Н	Н	⊣	٦	٦	⊣	႕	Н	, 	Н	٦	Н	Н	Н
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
394	σ	396	397	398	399	400	401	402	403	404	405	406	407	408	409	410	411	412	413	414

	222	0 0 0	222	777	TTGTCAAAGCCCTGGACGA TGTCAAAGCCCTGGACGAT STCAAAGCCCTGGACGATC
1 2	0 0		2 2	2 2	GTCAAAGCCCTGGACGATCTG TCAAAGCCCTGGACGATCTGC
1 2	2		7	7	CAAAGCCCTGGACGATCTGCA
1 2	7		2	2	AAAGCCCTGGACGATCTGCAC
1 2	7		2	2	AAGCCCTGGACGATCTGCACA
1 2	7		2	7	AGCCCTGGACGATCTGCACAA
1 2	7		7	7	GCCCTGGACGATCTGCACAAG
1 2	7		2	7	CCCTGGACGATCTGCACAAGA
1 2	7		2	2	CCTGGACGATCTGCACAGAT
1 2	7		7	7	CTGGACGATCTGCACAAGATG
1 2	7		7	7	TGGACGATCTGCACAAGATGA
1 2	7		7	2	GGACGATCTGCACAAGATGAA
1 2	7		7	7	GACGATCTGCACAAGATGAAC
1 2	7		7	7	ACGATCTGCACAAGATGAACC
1 2	7		7	2	CGATCTGCACAAGATGAACCA
1 2	2		2	7	GATCTGCACAAGATGAACCAC
1 2	2		2	7	ATCTGCACAGATGAACCACG
1 2	7		2	2	TCTGCACAGATGAACCACGT
2	2		7	7	CTGCACAGATGAACCACGTG

FIG. 20 (22)

2 2 2 2	1 2 2 2 GCACAAGATGAACCACGTGAC	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2	1 2 2 2
1	21	٦	J	J	J	7	٦	J	J	7	7	٦	7			႕	7		7	
437 2	438 2	439 2	440 2	441 2	442 2	443 2	444 2	445 2	446 2	447 2	448 2	449 2	450 2	451 2	452 2	453 2	454 2	455 2	456 2	457 2

FIG. 20 (23)

ACCCCCAACGTGT	GTCCCTGGG	CCCAACG	CCAACGTGTCCCTGGGCG	AACGTGTCCCTGGGCGC	GCTA	CTGGGCGCTAC			のいいでは、これのでは、これでは、これでは、これでは、これでは、これでは、これでは、これでは、これ	りかいいはまいかいのののでもこことにはいいかられているというというできます。	りりりつびょうのうのののこととしているというというのできません。	なてつりつりのでしていて	うらのでは、これでは、これでは、これでは、これでは、これでは、これでは、これでは、これ	つりつりのでしてい	CCTGGGCGCTACCGGGGGGCC	Ũ			つつつののののののののののののののののののののののののののののののののののの	GGCGCTACCGGGGGGGCCCCCG
220	2 0	3 6	~	7	Н	٦	7	Н		-	-	· -	, <u> </u>	٠,	\dashv	\vdash	\vdash			-
000	2 0	1 0	2	2	IJ	7	Н	7	г -1	러	·	l (-	· (-		-	7	٦	-	۱,	⊣
000	7 ~	1 7	2	7	႕	٦	-	Н	7	⊣	۲	-	_	1 -	-	⊣	7	႕	,	-
200	v ~	2	2	2	Н	႕	~	႕	Н	T		7	1	۱,	٠ ١		~ -1	1	_	4
777	v (2	. ~	٢	٦	-		Н	٦	٦	~	Н	۲	Н	-	⊣ •		, 	٦	_	4
21	21	21	21	21	21	21	21	21	21	21	21	21	21	10	J ,	7 7 7	21	21	21	1
50 60 60		62	63	64	65	99	29	89	69	70	71	72	73	7.4		7.7		77	28)

FIG. 20 (24)

GCGCTACCGGGGGGCCCCCGG	CGCTACCGGGGGGCCCCCGGC	GCTACCGGGGGGCCCCCGGCT	CTACCGGGGGGCCCCCGGCTG	TACCGGGGGGCCCCCGGCTGG	ACCGGGGGGCCCCCGGCTGGG	CCGGGGGCCCCCGGCTGGGC	CGGGGGCCCCCGGCTGGGGCC	GGGGGCCCCCGGCTGGGCCC	GGGGCCCCCGGCTGGGCCCG	GGGCCCCCGGCTGGGCCCGG	GGCCCCCGGCTGGGCCCGGG	GGCCCCCGGCTGGGCCCCGGGG	GCCCCGGCTGGGCCCCGGGGG	CCCCCGGCTGGGCCCCGGGGGC	CCCCGGCTGGGCCCCGGGGGCG	CCCGGCTGGGCCCCGGGGGCGT	CCGGCTGGGCCCCGGGGGCGTC	CGGCTGGGCCCCGGGGGGCGTCT	GGCTGGGCCCGGGGGCGTCTA	GCTGGGCCCGGGGGCGTCTAC	CTGGGCCCGGGGGCGTCTACG
٦	٦	\vdash	Н	\vdash	J	J	~-1	_	႕	_	- -	٦	 1	\vdash	J	 1	J	⊣	٦	~	_
٦	Н	Н	Н	٦	٠.	7	Н	П	႕	Н	Н	Н	႕	Н	~	٦	Н	٦	٦	٦.	IJ
٦	Н	٦	Н	Н	Н	٦	J	ᅼ	٦	Н	٦	٦	Н	Н	۲	٦	Н	۲	٦	٦	7
٦	H	٦	Н	Н	Н	Н	٦	Н	٦	Н	Н	Ч	Н	٦	Н	Н	H	٦	J	Н	٢
Н	Н	Н	 1	Н	Н	႕	Н	٦	႕	٦	Н	7	Н	٦	႕	႕	႕	Н	Н	Н	Н
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
62	081	181	182	183	184	185	981	187	88	681	061	191	192			195		197	198	499	200

FIG. 20 (25)

TGGGCCCGGGGGCGTCTACGC	CTACGCC	ACGCCG	CGCCGG	CCGGC	೧೦೮೮೧೦	CGGCCC	CGGCCCG	990009	GGCCCGGA	GCCCGGAG	CCCGGAGC	CGCCGGCCCGGAGCC	GAGCCA	AGCCAC	GCCACC	CCACCT	S	CCACCTCC	CCTCCC	CTCCCG
SGCCT	CGT	GGCCCGGGGGGCGTCTA	CCCGGGGGCGTCTA	STCTACG	CGTCTACG	GGGGCGTCTACGCCGGC	CGC	CGTCTACGCCGG	CGCCGG	CGCCGGC	CCGGCC	SGCCC	TACGCCGGCCCGGAGC	SCCGGAGCC	CCCGGAGCCA	CCGGCCCGGAGCCA	CGCCGGCCGGAGCCA	CGGAGCC	GC	AGCCAC
SCCGGC	5555555555	300000	366666	CCGGGGGCGTCTA	CGGGGGCGI	GGCGT	GGGCGTCTA	CGTCT	CGTCTA	CGTCTAC	TCTACG	CTACGC	ACGCCC	CGCCGGC	CGCCGG	CCGGCC	מפפככנ	CGGCCCC	GCCCGGA	೨೨၁၁၁
TGGG	0999	CCC	CCC	0000	SSSS	CGGG	9999	9999	0999	0000	GCGI	CGTC	GTCI	TCTA	CTAC	TACG	ACGC	CGCC	BCCG	5522
٦	Н	Н	Н	Н	Н	Н	Н	٦	Н	۲	Н	Н	Н	٦	Н	Н	٦	۲	۲	Н
Н	~	~	-	H	٦	, H	-1	7	Н	٦	7	-1	-	-	-	ᠳ	Н	Н	٦	٦
7	႕	7	٦	Ч	٦	٦	٦	٦	႕	٦	႕	7	٦	⊣	Н	1	٦	٦	٦	٦
7	٦	7	⊣	7	Н	۲	⊣	~	Н	Н	႕	⊣	⊣	⊣	rl	Н	Н	⊣	٦	Н
ч	٦	٦	Н	٦	٦	٦	⊣	Н	Н	٦	Н	Н	٦	Н	٦	7	Н	۲	Н	г - I
		21																		21
501	0		0	505	0	0	0	0	П	511	Ч	٦	٦		Н	\vdash	\vdash			521

FIG. 20 (26)

CGGCCCGGAGCCACCTCCCGT	GGCCCGGAGCCACCTCCCGTT	GCCCGGAGCCACCTCCCGT	CCCGGAGCCAC	CCGGAGCCACCTCCCGI	CGGAGCCACCTCCCGTTTAC	GGAGCCACCTCCCGTTTACAC	GAGCCACCTCCCGTTTACACC	AGCCACCTCCCGTTTACACCA	GCCACCTCCCGTTTACACC	CCACCTCCCGTTTACACCA	CACCTCCCGTTTAC	ACCTCCGGTTTACACCAACCT	CCTCCCGTTTACACCAACCTC	CTCCCGTTTACACCAACCTCA	TCCCGTTTACACCAACCTCAG	CCCGTTTACACCAACCTCAGC	CCGTTTACACCAACCTCAGC	CGTTTACACCAACCTCAGC	GTTTACACCAACCTCAGCAG	TTTACACCAACCTCAGCAGCT
Н	٦	7	٦	٦	Н	Н	٢	٢	Н	Ţ	۲	٦	7	2	7	2	7	7	~	٦
Н	Н	٦	1	٦	7	Н	7	7	٦	٦	Н	Н	7	2	7	2	2	7	-	ᠬ
٦	~	Н	Н	Н	щ	_		Ч	۲	Н	⊣	_	~	2	2	~	2	2	Н	-
7	٦	7	7	7	٦	Н	႕	٦	7	H	٦	Н	7	2	2	2	2	2	٦	Н
J	7	Н	Н	_	\vdash	٦		\vdash	7	_	\vdash	٦	Н	\vdash	\vdash	\vdash	Ч	_	\vdash	٦
				_	_	یے	_	_		ب	_				بے	ـــ			~	
		2]			2]						2]								2]	
522				2	N	528		\sim		\sim	\sim	\sim	ന		က	\sim	m			542

FIG. 20 (27)

TACACCAACCTCAGCAGCTA	ACACCAACCTCAGCAGCTAC	CACCAACCTCAGCAGCTAC	ACCAACCTCAGCAGCTACT	CCAACCTCAGCAGCTACTC	CAACCTCAGCAGCTACTCCC	AACCTCAGCAGCTACTCCCC	ACCTCAGCAGCTACTCCC	CCTCAGCAGCTACTCCCC	CCTCAGCAGCTACTCCCCAGC	CTCAGCAGCTACTCCCCAGCC	CAGCAGCTACTCCCCAGCCT	AGCAGCTACTCCCCAGCC	GCAGCTACTCCCCAGCCT	CAGCTACTCCCCAGCCTC	AGCTACTCCCCAGCCTCTGC	GCTACTCCCCAGCCTCTG	CTACTCCCCAGCCTCTGCGT	TACTCCCCAGCCTCTGCGTC	ACTCCCCAGCCTCTGCGTCC	CTCCCCAGCCTCTGCGTCCT	CTCCCCAGCCTCTGCGTCCTC
Н	Η	A	Ö	Ø	O	Ü	Ø	Ø	O	O	Ĥ	Ü	Ā	Ŋ	O	Ø	0	O	<u>-</u>	Ø	O
Н	~	2	7	Н	۲	Н	Ч	٦	Н	Н	Н	Н	Ч	Ч	Н	Н	Ч	Н	٦	۲	٦
Н	2	2	2	Н	Н	, 	 1	~	٦	Н	ᠳ	Н	~	, 	႕		-	٦	٦	7	Н
Н	2	2	2	Н	٦	Н	Ч	⊣	Н	Н	٦	Н	Н	Н	٦	٦	Н	Н	٦	٦	٦
Н	7	2	2	Н	٦	Н	~	Н	Н	႕	٦	Ч	٦	Н	۲	⊣	٦	٦	7	⊣	Н
Н	Н	⊣	Н	Ч	٦	Ч	Н	ᠬ	Н	Н	Н	Ч	Н	Н	Н	Н	Н	~	٦	٦	Н
					21																
543	-	4	4	4	548	4	5	\mathbf{c}	5	Ŋ	Ŋ	\mathbf{c}	\mathbf{S}	\mathcal{L}	$\mathbf{\sigma}$	\mathbf{c}	9	9	9	9	564

FIG. 20 (28)

CCCCAGCCTCTGCGTCCTCG	Ü	CCAGCCTCTGCGTCCTCGG	CAGCCTCTGCGTCCTCGGG	AGCCTCTGCGTC	GCCTCTGCGTCCTCGGGAG	CCTCTGCGTCCT	CTCTGCGTCCTCGGGAGGCG	CTCTGCGTCCTCGGGAGGCGC	TGCGTCCTCGGGAGGCGC	TGCGTCCTCGGGAGGCGC	GCGTCCTCGGG	CGTCCTCGGGA	STCCTCGGGAGGC	TCCTCGGGAGGC	CCTCGGGAGGCGCCGGGGC	CTCGGGAGGCGCCGGGGCTG		GGGAGGCGCCGGGGGGTTC	GGAGGCGCCGGGGCTGC	SGCGCGGGGCTGCC
1 T	ا د	1 C	1 C	1 C.	1 A	J G	J C			Ö	H	L G	ŭ	Ŋ] T	1 C	O	E	O	Ŋ
			-		•															
Н	႕	H	7	٦	Н	Н	Н	7	٦	۲	Н	Н	٦	Н	႕	Н	٦	٦	٦	Н
Н	Н	٦	Н	Н	 1	٦	⊣	Н	~	⊣	Н	٦	٦	႕	٦	Ч	٦	٦	٦	٦
Н	Н	Н	Ч	۲	Н	٦	႕	Н	٦	٦	٦	۲	ᆸ	٦	٦	Н	٦	Н	Н	٦
21	21	21	21	21		21				21										
- 1	266		568		7		/	7	7		7	1	7	7	∞	∞	∞	∞		585

586	21	٦	7	Н	Н	J	GGAGGCCCGGGGCTGCCGTC
37	21	-	Н	7	J	_	GAGGCGCCGGGGCTGCCGTCG
88		႕	٦	7	٦	_	AGGCGCCGGGCTGCCGTCGG
89		\vdash	Ч	~	H	٦	GGCGCCGGGCTGCCGTCGGG
		Н	٦	٦	٦	٦	GCGCCGGGGCTGCCGTCGGGA
91		Н	٦	٦	L	~	CGCCGGGGCTGCCGTCGGGAC
		Н	Н	~	٦	ᠳ	GCCGGGGCTGCCGTCGGGACC
93		Н	Ч	⊣	႕	٦	CCGGGGCTGCCGTCGGGACCG
94		٦	٦	٦	႕	႕	CGGGGCTGCCGTCGGGACCGG
95		~	Н	Н	႕	⊣	GGGGCTGCCGTCGGGACCGGG
		Ч	~	٦	٦	Н	GGGCTGCCGTCGGGACCGGGA
97	21	⊣	٦	٦	٦	Ч	GGCTGCCGTCGGGACCGGGAG
98		, -	٦	႕	7	٦	GCTGCCGTCGGGACCGGGAGC
66	21	٦	~	٦	٦	႕	CTGCCGTCGGGACCGGGAGCT
	21	⊣	Н	7	_	\dashv	TGCCGTCGGGACCGGGAGCTC
01	21	- -1	႕	⊣	ᆸ	٦	GCCGTCGGGACCGGGAGCTCG
02	21	~	Ч	-	٦	~	CCGTCGGGACCGGGAGCTCGT
03	21	٦	근	-	~	۲	CGTCGGGACCGGGAGCTCGTA
04	21	٦	~	٦	٦	႕	GTCGGGACCGGGAGCTCGTAC
05	21	٦	႕	٦	٦	٦	TCGGGACCGGGAGCTCGTACC
90	21	Н	۲	Н	J	IJ	CGGGACCGGGAGCTCGTACCC

FIG. 20 (30)

GGGACCGGGAGCTCGTACCCG	GGACCGGGAGCTCGTACCCGA	GACCGGGAGCTCGTACCCGAC	ACCGGGAGCTCGTACCCGACG	CCGGGAGCTCGTACCCGACGA	CGGGAGCTCGTACCCGACGAC	GGGAGCTCGTACCCGACGACC	GGAGCTCGTACCCGACGACCA	GAGCTCGTACCCGACGACCAC	AGCTCGTACCCGACGACCACC	GCTCGTACCCGACGACCACCA	CTCGTACCCGACGACCACCAT	TCGTACCCGACGACCACCATC	CGTACCCGACGACCACCATCA	GTACCCGACGACCACCATCAG	TACCCGACGACCACCATCAGC	ACCCGACGACCACCATCAGCT	CCCGACGACCACCATCAGCTA	CCGACGACCACCATCAGCTAC	CGACGACCACCATCAGCTACC	GACGACCACCATCAGCTACCT	ACGACCACCATCAGCTACCTC
7	٦	٦	٦	-	٦	٦	~	Н	٦	٦	٦	٦	٦	Н	7	7	~	7	7	7	2
7	Н	Н	ч	Н	Н	Н	Ч	Ч	٦	Н	Н	Ч	٦	Ч	7	7	2	2	7	2	2
Н	Н	Н	ㄷ	Н	Н	Н	٦	Н	⊣	Н	⊣	Н	⊣	ᆫ	7	2	2	2	2	7	2
1	٦	٦	٦	Н	۲	Н	Н	Н	٦	Н	٦	٦	٦	٦	2	2	2	7	7	7	2
٦	~	Ч	႕	٦	Н	⊣	႕	٦	႕	~	٦	⊣	٦	Н	႕	⊣	႕	٦	٦	٦	٦
21			21																		21
607	608	-		611					\vdash	-			2		622	623	624	625	626	627	628

FIG. 20 (31)

CGACCACCATCAGCTACCTCC	Ũ	TCAGCTACCTCCC	GCTACCTCCCA	C	CTCCCAC	CCATCAGCTACCTCCCACACG	CATCAGCTACCTCCCACACGC	ATCAGCTACCTCCCACACGCG	TCAGCTACCTCCCACACGCGC	CCAC	C	GCTACCTCCCACACGCGCCGC	CTACCTCCCACACGCGCCGCC	()	ACCTCCCACACGCGCCGCCT	CCTCCCACACGCGCCGCCCTT	CTCCCACACGCGCCGCCTTC	TTC	CCTTCG	CCTTCGC
2	2	2	~	2	2	2	~	٢	۲	Н	J	7	٦	⊣	Н	Н	٦	Н	٦	\vdash
2	2	7	8	2	7	2	2		Н	Н	Н	Н	٦	Н	7	Н	-	⊣	ᆏ	⊣
7	7	2	7	2	7	2	2	٦	٦	Н	Н	Н	IJ	-	႕	Н	٦	Н	Н	Н
2	7	7	2	2	7	2	7	Н	⊣	Н	⊣	Н	႕	~	~	Н	, 	⊣	Н	٦
٦	Н	Н	7	7	Н	ᅥ	~	7	٢	٦	ᆏ	러	႕	٦	٦	٦	Н	٦	٦	Н
21						21				21			21				21		21	21
59	30	31	32	33	34		36	37	38	39	40	41	42	43	44	45	46	47	48	49
9	છ	છે	છે	6	છે				છે	છે						9	9	9	79	79

FIG. 20 (32)

CACACGCCGCCCTTCGCCG	ACACGCGCCCCTTCGCCGG	CACGCCCCCCTTCGCCGGT	ACGCCCCCCTTCGCCGGTG	CECECCECCTTCGCCGGTGG	GCGCCGCCCTTCGCCGGTGGC	CGCCGCCTTCGCCGGTGGCC	GCCGCCCTTCGCCGGTGGCCA	CCGCCCTTCGCCGGTGGCCAC	CGCCCTTCGCCGGTGGCCACC	GCCCTTCGCCGGTGGCCACCC	CCCTTCGCCGGTGGCCACCCG	CCTTCGCCGGTGGCCACCCGG	CTTCGCCGGTGGCCACCCGGC	TTCGCCGGTGGCCACCCGGCG	TCGCCGGTGGCCACCCGGCGC	CGCCGGTGGCCACCCGGCGCA	GCCGGTGGCCACCCGGCGCAG	CCGGTGGCCACCCGGCGCAGC	CGGTGGCCACCCGGCGCAGCT	GGTGGCCACCCGGCGCAGCTG
٦	٦	⊣	Н	۲	٦	⊢ -	٦	۲	٦	Н	⊣	Н	႕	Н	⊣	Н	Н	Н	~	٦
7	1	٦	Н	۲	٦	Н	Ч	Н	-1	٦	-	Н	-	-	Н	٦	~	⊣	٦	⊣
~	ㄷ	Н	⊣	٦	႕	٦	Н	٦	٦	Ч	Н	П	႕	٦	႕	႕	۲	Т	1	1
٦	Ч	٦	٦	Ч	႕	Н	Н	Н	 1	Н	Н	Ч	٦	Н	Н	⊣	Н	٦	٦	Н
7	1	٦	٦	٦	٦	⊣	-	Н	Н	٦	٦	٦	IJ	٦	٦	1	Н	Н	Н	۲
21		21	21		21	21	21						21		21	21	21	21	21	21
വ	S	S				929					661				999			899		019

FIG. 20 (33)

GIGGCCACCCGGCGCAGCIGG	TGGCCACCCGGCGCAGCTGGG	GGCCACCCGGCGCAGCTGGGC	GCCACCCGGCGCAGCTGGGCT	CCACCCGCCCAGCTGGGCTT	CACCCGCCCAGCTGGGCTTG	ACCCGCCCAGCTGGGCTTGG	CCCGGCGCAGCTGGGCTTGGG	CCGCCCAGCTGGGCTTGGGC	CGGCGCAGCTGGGCTTGGGCC	GGCGCAGCTGGGCTTGGGCCG	GCGCAGCTGGGCTTGGGCCGC	CGCAGCTGGGCTTGGGCCGCG	GCAGCTGGGCCTTGGGCCGCGG	CAGCTGGGCTTGGGCCGCGGC	AGCTGGGCTTGGGCCGCGCG	GCTGGGCTTGGGCCGCGCGCGC	CTGGGCTTGGGCCGCGCGCC	TGGGCTTGGGCCGCGCCCT	GGGCTTGGGCCGCGCGCCTC	GGCTTGGGCCGCGCGCCTCC	GCTTGGGCCGCGCCTCCA
Ч	٦	٦	\vdash	٦	Н	Н	Н	Н	Н	٦	႕	٦	7	٦	Н	٦	٦	Н	Н	Н	۲
																				H	H
Н	~	⊣	۲	-	7	7	J	Ţ	J	۲	-	٦	٢	Н		٦	٦	٦	٦	٦	7
1	٦	Ч	Н	Н	~	Н	⊣	႕	Н	Н	٦	⊣	⊣		-	1	1	۲	۲	۲	Н
٦	Н	Н	Н	Н	Н	Н	Н	٦	٦	٦	~	٦	Н	Н	٦	٦	٦	٦	۲	٦	Н
21	21			21			21		21						21					21	21
7	572	~	1		7	~	1			∞							888		069	691	692
9	9	Ψ)	9	9	9	9	9	W	9	9	9	9	9	9	9	ဖ	9	9	9	J	4)

FIG. 20 (34)

でなり出してないかかったしつかかが出出し	ンダンン「ソンのつののつのつのつのではしていていている」	ばいい Fいいのいののいのいのいのいのい	ないではなっている。	ない こうりゅうりゅうしゅつ できる はっちょう しょうしょう しょうしょうしょう しょうしょう しょうしょう しょうしょう しょうしょう しょうしょう しょうしょう しょうしょう しょうしょう しょうしょうしょう しょうしょう しょうしょうしょう しょうしょうしょう しょうしょう しょう	CONTRACTOR OF THE CONTRACTOR O))T))のうちゅうちつう	せいしていりのうのからい	せいしていりのかのから	GCGGCGCCICCACCIICAAGG	CGCCCTCCACCTTCAAGGA	GGCGCCTCCACCTTCAAGGAG	GCGCCTCCACCTTCAAGGAAC	CACC		して正し	してくして	ACCLL	TCCACCTTCAAGGAGGAACCG	CCACCTTCAAGGAGGAAGAA		; (ACCITCAAGGAGGAACCGCAG	CCTTCAAGGAGGAACCGCAGA
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Н	1	Н	-	~	┍┥	· H	~	7	· -	٠,	1	~	٦	۲	٦	7	-، ا	-	, 	щ	_	Η,	1
21	21	21	21	21	21	21	21	21	2.1	1 6	T 7	21	21	21	21	21							
693	694	695	969	697	869	669	700	701	0	•	\supset	0			707	708	Č) (Η	711	٦	٦ ا	

FIG. 20 (35)

CTTCAAGGAGGAACCGCAGAC	TTCAAGGAGGAACCGCAGACC	TCAAGGAGGAACCGCAGACCG		AAGGAGGAACCGCAGACCGTG	AGGAGGAACCGCAGACCGTGC	GGAGGAACCGCAGACCGTGCC	GAGGAACCGCAGACCGTGCCG	(7)	GGAACCGCAGACCGTGCCGGA	GAACCGCAGACCGTGCCGGAG	U	ACCGCAGACCGTGCCGGAGGC	CCGCAGACCGTGCCGGAGGCG	CGCAGACCGTGCCGGAGGCGC	GCAGACCGTGCCGGAGGCGCG		AGACCGTGCCGGAGGCGCGCA	GACCGTGCCGGAGGCGCGCA	STGCCGGAGGCGCGC	GTGCCGGAGGCGCGCAGC
~	Н	Н	Н	Н	Н	٦	7	7	7	m	7	7	٦	٦	٦	Ч	н	-	ત	٦
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٦	Н	-	٦	Н	٦	Н	~	7	2	m	7	7	Н	┍╌┤	႕	\leftarrow	٦	Н	٦	7
Н	7	Ч	႕	Н	_	7	2	7	7	က	2	2	⊣	~	 1	Н	러	⊣	٦	۲
٦	~	⊣	٦	7	-	_	٦	٦	⊣	۲	٦	~	7	٦	Н	~	Н	٦	Н	Т
21	21	21	21	21	21	21	21	21	21	21	21	21		21	21	21	21	21	21	21
714		716	717	\vdash			2			724		7	2		0	730	731	732	733	734

FIG. 20 (36)

GTGCCGGAGGCGCGCAGCCGGGCGCGGGGGGGGGGGGGCGCGCGCGCGGGGGG			പപപപപപപപപപവ 2 2 2 2 2 2 2 2 2 2 2 2 2 2		
GGACGCCACGCCGGTG	ı — -	i -		ı — -	
GG	Н	ᠳ	IJ	-	
CGGGACGCCACGCCGCGG	۲	Н	٦	IJ	
CCGGGACGCCACGCCGCCG	Т	႕	Н	Н	
GCCGGGACGCCACGCCGCC	Н	 -	Н	٦	
AGCCGGGACGCCACGCCGC	2	2	7	7	
CAGCCGGGACGCCACGCCG	7	7	7	2	
GCAGCCGGGACGCCACGCC	7	7	7	7	
CGCAGCCGGGACGCCACGC	۲	Н	٦	П	
GCGCAGCCGGGACGCCACG	۲	~	근	IJ	
CGCGCAGCCGGGACGCCAC	٦	٦	Н	Н	
GCGCGCAGCCGGGACGCCA	٦	⊣	-	~	
GGCGCGCAGCCGGGACGC	Т	۲	႕	-	
AGGCGCGCAGCCGGGACG	႕	 1	႕	-	
SAGGCGCGCAGCCGGGACG	Н	႕	ᆏ	٦	
GGAGGCGCGCAGCCGGGAC	Ч	ᆏ	႕	٦	
CGGAGGCGCGCAGCCGGGA	Н	~	←I	Н	
CCGGAGG	Н	Н	щ	Н	
GCCGGAGGCGCGCAGC	Н	႕	Н	٦	
TGC	⊣	ч	Н	ᆸ	
ΤG	⊣	 1	Н	Н	

FIG. 20 (37)

GACGCCACGCCGGTGTCC	ACGCCACGCCGCCGGTGTCCC	CGCCACGCCGCCGGTGTCCCC	GCCACGCCGCCGGTGTCCCCC	CCACGCCGGGTGTCCCCCA	CACGCCGCCGGTGTCCCCCAT	ACGCCGCCGGTGTCCCCCCATC	CGCCGCCGGTGTCCCCCCATCA	GCCGCCGGTGTCCCCCCATCAA	CCGCCGGTGTCCCCCATCAAC	CGCCGGTGTCCCCCATCAACA	GCCGGTGTCCCCCATCAACAT	CCGGTGTCCCCCATCAACATG	CGGTGTCCCCCATCAACATGG	GGTGTCCCCCATCAACATGGA	GTGTCCCCCATCAACATGGAA	TGTCCCCCATCAACATGGAAG	GTCCCCCATCAACATGGAAGA	TCCCCCATCAACATGGAAGAC	CCCCCATCAACATGGAAGACC	CCCCATCAACATGGAAGACCA
~	7	~	2	7	2	2	2	2	7	~	7	2	7	7	7	7	2	7	7	7
7	2	2	2	2	2	2	2	7	2	~	2	2	2	2	2	7	. 7	2	2	2
2	7	7	7	7	2	7	7	7	7	7	7	7	2	7	2	7	~	7	7	2
2	2	7	2	7	7	2	7	0	7	7	7	7	2	7	2	7	2	7	7	2
Ä	Н	ᠬ	٦	Н	٦	7	1	Н	ᠳ	႕	٦	⊣	ᠬ	Н	2	2	7	2	2	2
21	21	21	21		21	21	21	21	21	21	21		21	21	21	21	21	21	21	21
Ŋ	758	759		9	762			9	9	167	9	9	7	771	7	773		775	116	111

FIG. 20 (38)

8 21 1 2 2 CCATCAACATGGAAGACCAAG 9 21 1 2 2 CATCAACATGGAAGACCAAG 1 21 1 2 2 2 CATCAACATGGAAGACCAAGA 2 21 1 2 2 2 ATCAACATGGAAGACCAAGA 3 21 1 2 2 2 AACATGGAAGACCAAGAGCGAAGAGCGCAAGAGCGCAAGAGCGCAAGAGCGCAAGAGCCCAAGAGCCCAAGAGCCCAAGAGCCCAAGAGCCCAAGAGCCCAAGAGCCCAAGAGCCCAAGAGCCCAAGAGCCCAAGAGCCCAACAGAGCCCATCAAAGAGCCCATCAAAGAGCCCAAGAGCCCATCAAAAGAGCCCATCAAAAGAGCCCATCAAAAGAGCCCATCAAAAGAGCCCAAGAGCCCATCAAAAGAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAGTGGAAGCCCATCAAAAAGTCAAAAAGTCAAAAAGTCAAAAAGTCAAAAAGTCAAAAAGTCAAAAAGTCAAAAAAAA	A	ָט ב	Ą	Ŋ	ü	ည	ပ္ပ	CA	Ţ	ຸບ	CA	Ą	A	Ŋ	Ħ	ט	Ŋ	Ą	ט	Ö	
8 21 1 21 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	ACC.	CCA	CAAC	AAG?	AGAC	GAG	AGC	SCGC	CGCZ	GCA	AT	ATC	TCA	CAA	AAA	AAG	AGTO	GTG	TGG	GGAG	して ペンプロ
8 21 1 21 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	CAAC	AAGA	AGAC	GACC	ACCA	CCAA	CAAG	AAGA	AGAG	GAGC	AGCG	CCCC	CGCA	GCAT	CATC	ATCA	TCAA	CAAA	AAAG	AAGT	じたして
8 21 1 21 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	ATG	4TGG	rgga	GAA	SAAG	AAGA	AGAC	SACC	ACCA	CCAA	CAAG	AAGA	AGAG	SAGC	AGCG	SCGC	CGCA	SCAT	CATC	ATCA	じょくないじ
8 21 1 21 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	AAD	'AAC	ACA	CAT	ATG(TGG/	GGA	GAA	:AAG	AGA(GAC	ACC!	CCA	CAAC	:AAG	AGAC	GAGG	;AGC(GCGC	CCC	してしてしてる
8 21 1 21 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	ראטטר	CCAT	CATCA	ATCA?	PCAAC	CAAC	AACAT	ACATO	CATGG	ATGG?	rgga2	SGAAG	SAAGE	AAGAC	AGACC	SACCA	ACCAA	CCAAG	SAAGA	AAGAG	שלשלע
8 21	Ī														-						2
8 21																					
8 8 2 1 1 2 1 1 2 2 2 2 1 1 1 2 3 2 1 1 1 1	~	2	7	7	7	7	7	7	7	7	2	7	7	7	2	7	7	~	7	2	^
88 21 0 0 8 4 3 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	2	0	7	7	7	7	7	2	2	2	7	7	2	7	2	2	2	2	2	2	^
88 21 0 0 8 4 3 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	0	~ 1	~1	~ 1	~ 1	01	01	01	01	01	01	01	01	01	01	~ 1	01	01	~ 1	01	•
8 4 3 2 4 3 2 1 0 9 8 4 6 5 4 3 2 1 1 1 1 1 1 2 2 2 2 2 2 1 1 1 1 1 1	•			•				.,				.,		17	(1	.,	.,	.,	()	.,	
80010645078001064507	7	٦	٦	⊣	H	Ч	Н	Н	Н	Н	Н	Н	ᅼ	Н	٦	Н	٦	Ч	٦	Н	_
80012645978001264597	21																	21			2.1
		0	08	81 ;	82 ;	83 2	4	_	86	87	œ		_	91 2				95.		97	98

FIG. 20 (39)

799	21	7	7	7	7	2	GAGCGCATCAAAGTGGAGCGC
800	21	7	2	7	7	7	AGCGCATCAAAGTGGAGCGCA
801	21	-1	7	7	2	2	GCGCATCAAAGTGGAGCGCAA
802	21	Ч	7	7	2	2	CGCATCAAAGTGGAGCGCAAG
803	21	П	7	7	2	2	GCATCAAAGTGGAGCGCAAGC
804	21	Н	7	7	2	2	CATCAAAGTGGAGCGCAAGCG
805	21	٦	7	7	2	7	ATCAAAGTGGAGCGCAAGCGG
806	21	Н	7	7	2	2	TCAAAGTGGAGCGCAAGCGGC
807	21	Т	7	7	2	7	CAAAGTGGAGCGCAAGCGGCT
808	21	႕	7	7	2	2	AAAGTGGAGCGCAAGCGGCTG
809	21	Н	7	2	2	7	AAGTGGAGCGCAAGCGGCTGC
810	21	7	7	7	7	7	AGTGGAGCGCAAGCGGCTGCG
811	21	႕	7	7	2	7	GTGGAGCGCAAGCGGCTGCGG
812	21	러	7	2	7	7	TGGAGCGCAAGCGGCTGCGGA
813	21	႕	7	7	7	2	GGAGCGCAAGCGGCTGCGGAA
814	21	Ч	7	7	7	2	GAGCGCAAGCGGCTGCGGAAC
815	21	႕	٦	٦	J	Н	AGCGCAAGCGGCTGCGGAACC
816	21	႕	7	٦	Н	Н	GCGCAAGCGGCTGCGGAACCG
817	21	٦	٦	٦	Н	Н	CGCAAGCGGCTGCGGAACCGG
818	21	٦	Н	٦	Н	Н	GCAAGCGGCTGCGGAACCGGC
819	21	Ч	႕	٦	Ч	႕	CAAGCGGCTGCGGAACCGGCT
820	21	٦	7	7	7	2	AAGCGGCTGCGGAACCGGCTG

AGCGGCTGCGGAACCGGCTGG		CGGCTGCGGAACCGGCTGGCG	GGCTGCGGAACCGGCTGGCGG	GCTGCGGAACCGGCTGGCGGC	CTGCGGAACCGGCTGGCGGCC	TGCGGAACCGGCTGGCGGCCA	GCGGAACCGGCTGGCGGCCAC	CGGAACCGGCTGGCGGCCACC	GGAACCGGCTGGCGGCCACCA	GAACCGGCTGGCGGCCACCAA	AACCGGCTGGCGGCCACCAAG	ACCGGCTGGCGGCCACCAAGT	CCGGCTGGCGGCCACCAAGTG	CGGCTGGCGGCCACCAAGTGC	GGCTGGCGGCCACCAAGTGCC	GCTGGCGGCCACCAAGTGCCG	CTGGCGGCCACCAAGTGCCGG	TGGCGGCCACCAAGTGCCGGA	GGCGGCCACCAAGTGCCGGAA	GCGCCACCAAGTGCCGGAAG
7	~	7	2	2	7	8	8	2	2	7	7	2	2	2	2	2	7	2	2	2
2	2	7	7	7	2	2	2	2	2	2	7	7	7	2	7	7	2	7	2	7
2	7	7	7	2	7	7	7	7	2	2	7	7	7	7	7	2	7	7	7	2
2	7	7	2	2	7	2	7	2	2	2	2	2	7	2	2	2	2	2	2	2
П	Н	٦	ᆏ	٦	٦	٦	Н	٦	٦	~	⊣	ᡤ	٦	П	2	7	7	7	7	7
21	21	21	21	21	21									21					21	21
821	822	823	824	825	826	-			830	831	832	833	834	835	836	837	838	839	840	841

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FIG. 20 (41)

CGGCCACCAAGTGCCGGAAGC	GGCCACCAAGTGCCGGAAGCG	GCCACCAAGTGCCGGAAGCGG	CCACCAAGTGCCGGAAGCGGA	CACCAAGTGCCGGAAGCGGAA	ACCAAGTGCCGGAAGCGGAAG	CCAAGTGCCGGAAGCGGAAGC	CAAGTGCCGGAAGCGGAAGCT	AAGTGCCGGAAGCGGAAGCTG	AGTGCCGGAAGCGGAAGCTGG	GTGCCGGAAGCGGAAGCTGGA	TGCCGGAAGCGGAAGCTGGAG	GCCGGAAGCGGAAGCTGGAGC	CCGGAAGCGGAAGCTGGAGCG	CGGAAGCGGAAGCTGGAGCGC	GGAAGCGGAAGCTGGAGCGCA	GAAGCGGAAGCTGGAGCGCAT	AAGCGGAAGCTGGAGCGCATC	AGCGGAAGCTGGAGCGCATCG	GCGGAAGCTGGAGCGCATCGC	CGGAAGCTGGAGCGCATCGCG
7	2	~	7	~	2	7	7	7	7	7	7	7	7	7	~	7	2	7	7	7
2	7	2	2	2	2	7	7	2	2	2	2	2	2	7	2	2	2	2	2	2
~	~	7	7	7	7	7	7	~	2	7	7	7	2	7	7	~	7	7	2	2
2	2	2	7	2	2	2	2	2	2	2	2	2	2	2	7	7	7	7	7	7
2	2	2	7	2	7	2	2	2	2	2	7	2	2	2	2	7	2	2	2	7
21	21	21			21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
842	843	844	845	846	847	848	849		851		853	854	855	856	857	858	859	_	861	862

FIG. 20 (42)

GCGCATCGCGC	U	CATCGC	CATCGCGCCC	ATCGCGCGCCT	TCGCGCGCCTG	SCGCGC	CGCGCC	CGCCCTGGAG	GCGCCTGGAGG	CGCC	CGCCTGGAGGAC	CCTGGAGGACA	CTGGAGGAC	ATCGCGCCCTGGAGGACAAG	CCTGGAGGACAAGG	TGGAGGACAAGGT	CGCCTGGAGGACAAGGTG	CGCGCCTGGAGGACAAGGTGA	GACAAGGTGAA	CCTGGAGGACAAGGTGAAG	CAAGGTGAAGA
GGAAGCTGGAGCGCATCG	GAAGCTGGAGCGCATCG	AAGCTGGAGCG	AGCTGGAGCGC	GCTGGAGCGCATCG	CTGGAGCGCA	TGGAGCGCATC	GGAGCGCATCG	GAGCGCATCG	AGCGCATCGCG	GCGCATCGCG	CGCATCGCGC	GCATCGCGCG	CATCGCGCGC	ATCGCGCGCC	TCGCGCGCCT	SCCCCCCTG	GCGCGCCTGG	CGCGCCTGGA	GCGCCTGGAGGA	CGCCTGGAGG	GCCTGGAGGA
7	7	7	2	7	7	7	2	2	7	7	7	7	7	7	7	7	7	2	7	7	7
2	2	2	2	7	7	7	7	7	2	2	7	2	2	7	2	2	7	2	2	2	2
7	7	7	7	7	7	7	7	7	7	2	7	7	7	7	7	7	7	7	2	2	2
2	7	7	7	7	7	7	2	7	2	2	7	2	7	7	7	7	7	7	2	7	2
2	2	7	7	7	7	2	7	7	2	7	7	7	7	7	7	7	7	7	2	2	2
21	21													21							21
863	864	865	998	867	868	869	7		872	~	-	875	876	-	878	879		881	882	883	884

FIG. 20 (43)

		TGGAGGACAAGGTGAAGACACC	GGAGGACAAGTAGTAAAAGAGGG	GAGGACAAGGTGAAGACGCI	AGGACAAGGTGAAGACGCT	GGACAAGGTGAAGACGCTCAA	GACAAGGTGAAGACGCTCAAG	ACAAGGTGAAGAAGACGTTAAAGG	CAAGGTGAAGAAGTGAACTGAACTGAACTGAACTGAACT	AAGGTGAAGACGCTCAAGGC	AGGTGAAGACTCACACACACACACACACACACACACACAC	GGTGAAGACGCTCAACGCCG	GTGAAGACGCTCAAGGCCGA	TGAAGACGCTCAAGGCCGAGA	GAAGACGCTCAAGGCCGAGAA	AAGACGCTCAAGGCCGAAAC	AGACGCTCA AGACGCCCCCCCCCCCCCCCCCCCCCCCCC	OD CACOLLOS A CALOS CONTRACTOR OF CALOS CONTRACTOR OF CALOS	CGCTCAAGG	GCTCAAGGC
~	ι (\)	2	~	2	2	2	2	2	2		~	· —		٦	٦	-	,		. —	
2	2	2	2	2	7	2	2	2	2	Н	~	· ~	H	-	~	-	_	_	l	Н
7	2	7	2	2	23	2	2	7	2	1	٦	~	Н	Н	Н	Н	~	-	7	Н
2	2	7	2	2	2	7	7	2	2	⊣	Н	Н	Н	٦	۲	٦	۲	Н	٦	П
2	Н	Н	ᆏ	-	Н	-	۲	Н	٦	Н	٦	٦	٦	٦	-	, 	러	⊣	۲	٦
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21		21	21
885	886		888	889	890	891		σ	894	895	9	Ó	868	g			902	903	904	902

FIG. 20 (44)

GCTCAAGGCCGAGAACGCGG	CICAAGGCCGAGAACGCGGG		AAAGCCGAGA AAGCCGG	うりつうつう Taring Control A Public Control A Published A	GCCGAGAACGCGGGGCTT	CGAGAACGCGGGCTGT	AACGCGGGGCTGTC	GAGAACGCGGGGCTGTCG	GAACGCGGGGGCTGTCG	AGAACGCGGGCTGTCGAG	GAACGCGGGGCTGTCGACT	GGCTGTCGAGTA	GCTGTCGAGTAC	TGTCGAGTACC	TACCG	TCGAGTACCGC	GGGCTGTCGAGTACCGC	TGTCGAGTACCCC	GCTGTCGAGTACCGCCG	TCGAGTACCGCCGGC
Н	~	~	· ~	2	~	7	2	2	2	2	2	7	-	٦	Ч	7		Н	٦	٦
٦	7	2	~	2	2	2	7	7	2	2	2	2	Н	٦	۲	Н	Н	П	~	Н
۲	2	7	2	2	7	7	2	~	7	7	7	7	٦	႕	۲	Н	Н	٦	Н	٦
7	2	7	2	2	2	7	2	2	2	2	2	2	٦		Н	Н	႕	Н	-	٦
٦	7	۲	Н	٦	H	⊣	Н	႕	٦	7	٦	Н	Н	Н	7	٦	۲	۲	٦	П
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
90	07	08	60	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	56
٥١	U١	U١	U	U١	U)	ΟΊ	Ų١	O1	ΟJ	0	Ω)	ΟJ	ΟJ	O١	σı	σ)	σ	σ	σ	σ

FIG. 20 (45)

GCTGTCGAGTACCGCCGGCCT	TCGAGTACCGCCGGCC	CCCCGGCCTC	CCTCC	TCGAGTACCGCCGGCCTCCTC	ľĊ	Õ	AGTACCGCCGGCCTCCTCCGG	CTCCG	TACCGCCGCCTCCTCCGGGA	ACCGCCGCCTCCTCCGGGAG	CCGCCGCCTCCTCCGGGAGC	CGCCGGCCTCCTCCGGGAGCA	GCCGGCCTCCTCCGGGAGCAG	CCGGCCTCCTCCGGGAGCAGG	CGGCCTCCTCCGGGAGCAGGT	GGCCTCCTCCGGGAGCAGGTG	GCCTCCTCCGGGAGCAGGTGG	Ü	\mathcal{O}	$C_{\mathcal{O}}$	CCTCCGGGAGCAGGTGGCCCA
Н	Н	Н	٦	Н	٦	Н	Н	Н	٦	٦	٦	٦	٦	Н	Н	٦	٦	Н	٦	٦	~
			٦																		- -
_	П		7	1		_	7	_	7			7	_		П			7	_	-	7
Н	Н	٦	Н	Н	~	Н	٦	-	٦	~	۲-	⊣	. 	Н	٦	⊣	⊣	٦	Н	Н	Н
Н	Н	٦	П	Н	٦	Н,	<u>,</u> l	7	٦	П	7	1	J	7	Н	Н	٦	۲	ᆏ	⊣	Н
21			21	21			21			21		21							21	21	21
927	_	_	930	-	\mathcal{C}				\sim		m	n	4		4		944		946	947	948

FIG. 20 (46)

TCCGGGAGCAGGTGGCCCAG	CCGGGAGCAGGTGGCCCAGC	CGGGAGCAGGTGGCCCAGCT	GGGAGCAGGTGGCCCCAGCTC	GGGAGCAGGTGGCCCAGCTCA	GGAGCAGGTGGCCCAGCTCAA	STGGCCCAGCTCAAA	AGCAGGTGGCCCAGCTCAAAC	SGCCCAGCTCAAACA	SCCCAGCTCAAACAG	CCCAGCTCAAACAGA	CCAGCTCAAACAGAA	CCCAGCTCAAACAGAAG	TGGCCCAGCTCAAACAGAAGG	CAGCTCAAACAGAAGGT	CAGCTCAAACAGAAGGTC	CAGCTCAAACAGAAGGTCA	CCAGCTCAAACAGAAGGTCAT	GCTCAAACAGAAGGTCATG	CTCAAACAGAAGGTCATGA	CAAACAGAAGGTCATGAC
CTCCGG	TCCGGG	CCGGGGA	CGGGAGG	GGGAGC	GGAGCA	GAGCAGGTGG	AGCAGG	GCAGGTG	CAGGTGG	AGGTGG	GGTGGCC	GTGGCC	TGGCCC?	GGCCCA	GCCCAG	CCCAGC	CCAGCT	CAGCTCA	AGCTCA	GCTCAA
Н	٦	۲	Н	7	۲	7	2	2	7	2	7	٦	J	⊣	Н	٦	Н	7	2	7
Н	٦	H	٦	٦	٦	7	2	2	2	2	7	႕	⊣	႕	٦	٦	٦	2	2	7
Н	Н	러	Н	Н	Н	2	2	2	2	2	2	⊣	Н	Н	⊣	۲	Н	2	2	7
7	~	Н	Ч	٦	٦	2	2	2	2	2	2	۲	~	-	٦	~	٦	2	2	2
٦	٦	Н	Н	Н	H	٦	٦	۲	Н	႕	ᠬ	Н	٦	Н	Н	٦	٦	٦	႕	Н
21													21				21		21	21
949				Ŋ	954	Ŋ		Ŋ				9	962	963	964	965	996	9	896	696

FIG. 20 (47)

CTCAAACAGAAGGTCATGACC	TCAAACAGAAGGTCATGACCC	CAAACAGAAGGTCATGACCCA	AAACAGAAGGTCATGACCCAC	AACAGAAGGTCATGACCCACG	ACAGAAGGTCATGACCCACGT	CAGAAGGTCATGACCCACGTC	AGAAGGTCATGACCCACGTCA	GAAGGTCATGACCCACGTCAG	AAGGTCATGACCCACGTCAGC	AGGTCATGACCCACGTCAGCA	GGTCATGACCCACGTCAGCAA	GTCATGACCCACGTCAGCAAC	TCATGACCCACGTCAGCAACG	CATGACCCACGTCAGCAACGG	ATGACCCACGTCAGCAACGGC	TGACCCACGTCAGCAACGGCT	GACCCACGTCAGCAACGGCTG	ACCCACGTCAGCAACGGCTGT	CCCACGTCAGCAACGGCTGTC	CCACGTCAGCAACGGCTGTCA
~	~	2	Н	Н	Н	2	2	2	2	~	~	~	~	~	2	7	7	٦	Н	~
		•				2 2														-
2	2	2	۲-	⊣	٦	2	2	2	2	2	2	7	7	7	2	2	7		٦	⊣
7	М	٦	٦	٦	٦	Н	-	٦	Н	Н	٦	٦	ᆏ	ㄷ	٦	٦	٦	٢	Н	٦
						21														
~	~		1	~	1	916	7		1	∞	∞	∞					ω			$\boldsymbol{\omega}$

FIG. 20 (48)

CACGTCAGCAACGGCTGTCAG	ACGTCAGCAACGGCTGTCAGC	⊢E⊣	CAGC	GCTG	CAGCTGC	AACGGCTGTCAGCTGCT	GCAACGGCTGTCAGCTGC	CAACGGCTGTCAGCTGCT	Ē	IGC	CGGCTGTCAGCTGCTTGG	CAGCTGCTGCTTG	GCTGTCAGCTGCTTGGGG	CTGTCAGCTGCTTGGGGGT	TGTCAGCTGCTGCTTGGGGTC	GTCAGCTGCTGCTTGGGGTCA	TCAGCTGCTGCTTGGGGTCAA	TTGGGGTCA	AGCTGCTTGGGGGTCAAGG	G	SGGGT
Ч	Н	~	Н	Н	Н	Ч	⊣	Н	<mark>근</mark>	۲	٦	٦	٦	۲	٦	٦	۲	Н	۲	٦	Н
٦	_	⊣	۲	П	Н	П	Ч	٦	٦	Н	-	٦	٦	Н	⊢	٦	Н	⊣	Н	⊣	Н
٦	1	႕	7	Н	1	7	႕	Н	٦	Н	Н	Н	Н	7	۲	Н	Н	Н	ㄷ	٦	٦
٦	-	Н	٦	ᆏ	ᆏ	⊣	Н	٦	~	ㄷ	Ч	႕	Н	Н	٦	⊣	႕	Ч	٦	Н	Н
7	٦	٦	۲	Н	⊣	Н	Н	٦	Н	٦	 1	Н	⊣	٦	러	~~i	⊣	Н	Н	٦	٦
						21											21			21	
9	992	9			-	9			1000	0	1002		1004	1005	1006	1007	1008	1009		1011	1012

	_					.	.		.		
CAAGGGAC	AAGGGACA	AGGGACAC	GGGACACG	GGACACGC	GACACGCC	ACACGCCT	CACGCCTI	ACGCCTTC	CGCCTTCT	GCCTTCTG	CCTTCTGA
TGCTGCTTGGGGTCAAGGGAC	TGCTTGGGGTCAAGGGACA	CTGCTTGGGGTCAAGGGACA	TGCTTGGGGTCAAGGGACACG	GCTTGGGGTCAAGGGACACGC	CTTGGGGTCAAGGGACACGCC	TTGGGGTCAAGGGACACGCCT	TGGGGTCAAGGGACACGCCTT	GGGGTCAAGGGACACGCCTTC	GGGTCAAGGGACACGCCTTCT	GGTCAAGGGACACGCCTTCTG	GTCAAGGGACACGCCTTCTGA
TGCTG	GCTGC	CIGCI	TGCTI	GCTTG	CTTGG	TTGGG	TGGGG	GGGGT	GGGTC	GGTCA	GTCAA
2	7	7	7	2	7	7	7	2	7	~	7
7	2	7	7	2	7	2	7	7	2	7	2
2	2	7	7	2	2	7	7	7	2	2	2
2	7	7	7	7	7	7	2	2	2	7	2
٦	٦	Н	Н	H	٦	-	Н	2	7	7	2
21	21	21	21	21	21	21	21	21	21	21	21
1013	1014	1015	1016	1017	1018	1019	1020	1021	1022	1023	1024

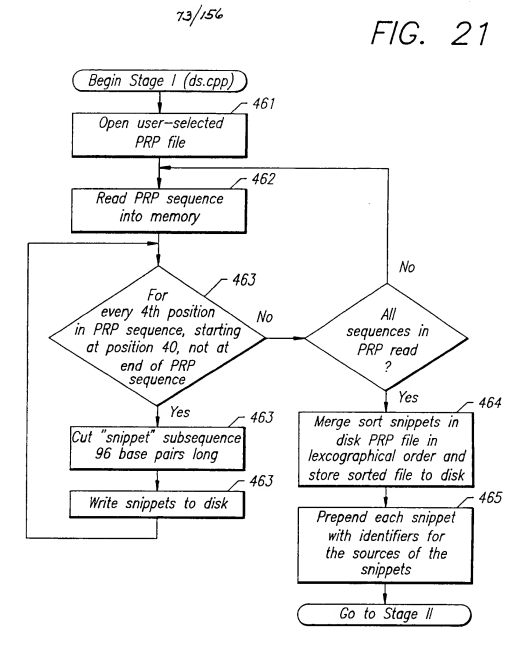
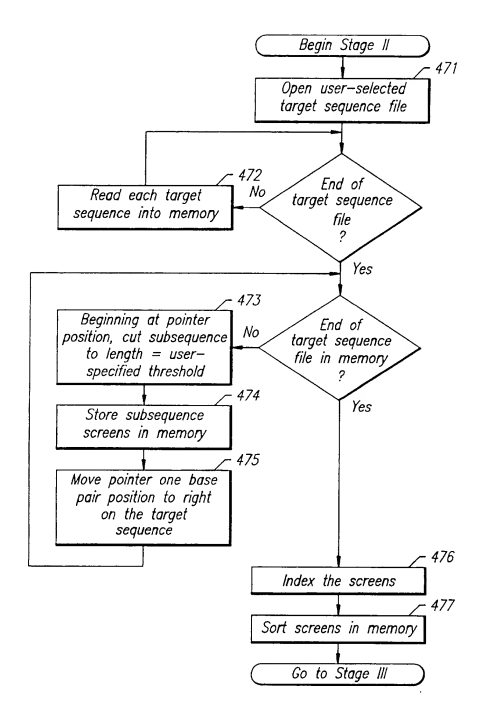
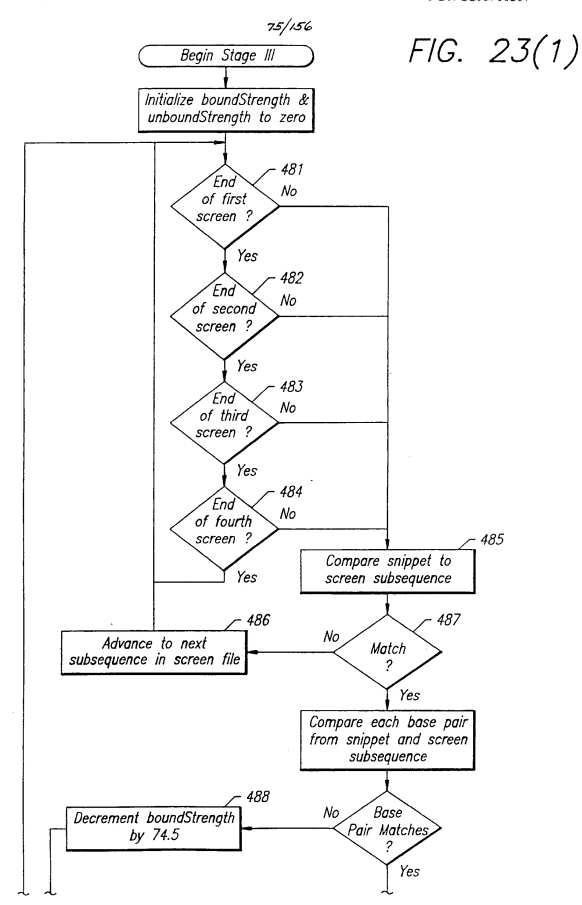


FIG. 22





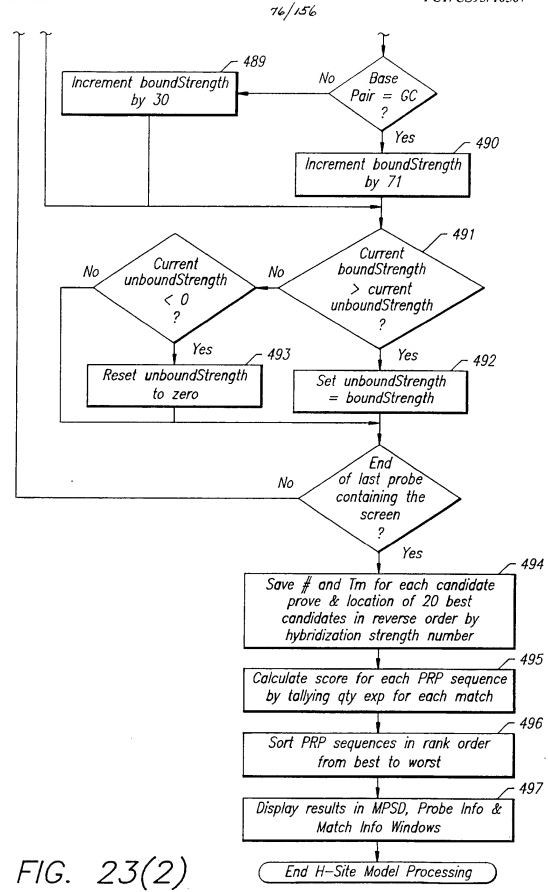


FIG. 24A (1)

OligoProbe DesignStation

Probes: C:\HITACHI\HUMBJUNX.CDS Datatbase: C:\HITACHI\JUNMIX.SEQ

= 4

11

Mismatch Model,

77/	156 0 Q										
ensN	Probe										
screensN	∞	ATGTGCACTAAAATGGAACAG	TGTGCACTAAAATGGAACAGC	GTGCACTAAAATGGAACAGCC	TGCACTAAAATGGAACAGCCC	GCACTAAAATGGAACAGCCCT	CACTAAAATGGAACAGCCCTT	ACTAAAATGGAACAGCCCTTC	CTAAAATGGAACAGCCCTTCT	TAAAATGGAACAGCCCTTCTA	AAAATGGAACAGCCCTTCTAC
	7	AAAATO	AAATG	AATGG	ATGGA	TGGAA	GGAAC	GAACA	AACAG	ACAGC	CAGCC
	9	GCACT	CACTA	ACTAA	CTAAA	TAAAA	AAAAT	AAAATG	AAATGG	ATGGA	TGGAA
	ហ	ATGI	\mathtt{TGTG}	GTGC	TGCA	GCAC	CACI	ACLA	CTAA	TAA	AAAA
	4	0	0	0	0	0	0	0	0	0	0
•	m	0	0	0	0	0	0	0	0	0	0
les	2	0	0	0	0	0	0	0	0	0	0
smatc	0 1 2	0	0	0	0	0	0	0	0	0	0
		0	0	0	0	0	0	0	0	0	0
ion	gth	21	21	21	21	21	21	21	21	21	21
Position	leng.	1	2	C	4	5	9	7	∞	6	10
•											

AAATGGAACAGCCCTTCTACC	AATGGAACAGCCCTTCTACCA	ATGGAACAGCCCTTCTACCAC	TGGAACAGCCCTTCTACCACG	GGAACAGCCCTTCTACCACGA	GAACAGCCCTTCTACCACGAC	AACAGCCCTTCTACCACGACG	ACAGCCCTTCTACCACGACGA	CAGCCCTTCTACCACGACGAC	AGCCCTTCTACCACGACGACT	GCCCTTCTACCACGACGACTC	CCCTTCTACCACGACGACTCA	~	CTTCTACCACGACGACTCATA	TTCTACCACGACGACTCATAC	TCTACCACGACGACTCATACA	CTACCACGACGACTCATACAC	TACCACGACGACTCATACACA	ACCACGACGACTCATACACAG	CCACGACGACTCATACACAGC	CACGACGACTCATACACAGCT
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
7	J	⊣	_	_		Ч	J	٦	Ч	Н		\vdash	J	\vdash	 1	7	\vdash	\vdash	7	7
~	7	7	7	2	7	7	2	2	~		2	7	7	~	8	7	~	2	~	7
11	12	13	14	15			18	19	20		22	23	24	25	26	27	28	29	30	31

(A)

ACGACGACTCATACACACA	GACGACTCATACACAGCTA	CACAGCTAC	ACGACTCATACACAGCTACGG	CGACTCATACACAGCTACGGG	GACTCATACACAGCTACGGGA	ACTCATACACAGCTACGGGAT	CTCATACACAGCTACGGGATA	TCATACACAGCTACGGGATAC	CATACACAGCTACGGGATACG	ATACACAGCTACGGGATACGG	TACACAGCTACGGGATACGGC	ACACAGCTACGGGATACGGCC	CACAGCTACGGGATACGGCCG	ACAGCTACGGGATACGGCCGG	CAGCTACGGGATACGGCCGGG	AGCTACGGGATACGGCCGGGC	GCTACGGGATACGGCCGGGCC	CTACGGGATACGGCCGGGCCC		ACGGGATACGGCCGGGCCCCT	CGGGATACGGCCGGGCCCCTG
С	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	Ō	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21		21	21		21		21	21	21	21	21	21	21	21	21	21	21
C)	~	4	2	9	7	ω	0	0	٦	2	43	4	വ	9	47	ω	σ	0	_	2	m

FIG. 24A (4)

						80	/15	-6												
GGGATACGGCCCGGGCCCCTGG	GGATACGGCCCGGGCCCCTGGT	GATACGGCCGGGCCCCTGGTG	ATACGGCCGGGCCCCTGGTGG	TACGGCCGGCCCCTGGTGGC	ACGCCCGGCCCCTGGTGGCC	CGGCCGGCCCCTGGTGGCCT	GGCCGGCCCCTGGTGGCCTC	GCCGGGCCCTGGTGGCCTCT	CCGGGCCCTGGTGGCCTCTC	CGGGCCCTGGTGGCCTCTCT	GGGCCCCTGGTGGCCTCTCTC	GGCCCCTGGTGGCCTCTCTCT	GCCCCTGGTGGCCTCTCTCTA	CCCCTGGTGGCCTCTCTAC	CCCTGGTGGCCTCTCTACA	CCTGGTGGCCTCTCTACAC	CTGGTGGCCTCTCTACACG	TGGTGGCCTCTCTACACGA	GGTGGCCTCTCTACACGAC	GTGGCCTCTCTCTACACGACT
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	Ö	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
54	52	26	57	28	59	09	61	62	63	64	65	99	29	68	69	70	71	72	73	74

FIG. 24A (5)

						8	1/1	56												
GGCCTCTCTACACGACT	GGCCTCTCTCTACACGACTAC	GCCTCTCTCTACACGACTACA	CCTCTCTACACGACTACAA	CTCTCTCTACACGACTACAAA	TCTCTCTACACGACTACAAAC	CTCTCTACACGACTACAAACT	TCTCTACACGACTACAAACTC	CTCTACACGACTACAAACTCC	TCTACACGACTACAAACTCCT	CTACACGACTACAAACTCCTG	TACACGACTACAAACTCCTGA	ACACGACTACAAACTCCTGAA	CACGACTACAAACTCCTGAAA	ACGACTACAAACTCCTGAAAC	CGACTACAAACTCCTGAAACC	GACTACAAACTCCTGAAACCG	ACTACAAACTCCTGAAACCGA	CTACAAACTCCTGAAACCGAG	TACAAACTCCTGAAACCGAGC	ACAAACTCCTGAAACCGAGCC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21																				
75	92	11	78	79	80	81	85	83	84	82	86	87	88	83	90	91	92	93	94	95

FIG. 24A (6)

							82,	/ 14	76										
CAAACTCCTGAAACCGAGCCT	AAACTCCTGAAACCGAGCCTG	AACTCCTGAAACCGAGCCTGG	ACTCCTGAAACCGAGCCTGGC	CTCCTGAAACCGAGCCTGGCG	TCCTGAAACCGAGCCTGGCGG	CCTGAAACCGAGCCTGGCGGT	CTGAAACCGAGCCTGGCGGTC	TGAAACCGAGCCTGGCGGTCA	GAAACCGAGCCTGGCGGTCAA	AAACCGAGCCTGGCGGTCAAC	AACCGAGCCTGGCGGTCAACC	ACCGAGCCTGGCGGTCAACCT	CCGAGCCTGGCGGTCAACCTG	CGAGCCTGGCGGTCAACCTGG	GAGCCTGGCGGTCAACCTGGC	AGCCTGGCGGTCAACCTGGCC		CTGGCC	CCGA
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
96	2	ω 8	σ	0	0	0	0	0	105	0	0	0	0	110	111	112	113	114	115

FIG. 24A (7)

TGGCGGTCAACCTGGCCGACC	GGCGGTCAACCTGGCCGACCC	GCGGTCAACCTGGCCGACCCC	CGGTCAACCTGGCCGACCCCT	GGTCAACCTGGCCGACCCCTA	GTCAACCTGGCCGACCCCTAC	TCAACCTGGCCGACCCCTACC	CAACCTGGCCGACCCCTACCG	AACCTGGCCGACCCCTACCGG	ACCTGGCCGACCCCTACCGGA	CCTGGCCGACCCCTACCGGAG	CTGGCCGACCCCTACCGGAGT	TGGCCGACCCCTACCGGAGTC	GGCCGACCCCTACCGGAGTCT	GCCGACCCCTACCGGAGTCTC	CCGACCCCTACCGGAGTCTCA	CGACCCCTACCGGAGTCTCAA	GACCCCTACCGGAGTCTCAAA	ACCCCTACCGGAGTCTCAAAG	CCCCTACCGGAGTCTCAAAGC	CCCTACCGGAGTCTCAAAGCG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	Ö	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
16	17	18	19	20	21	.22	23	24	.25	.26	.27	.28	53	30	31	.32	.33	34	35	36

FIG. 24A (8)

CCTACCGGAGTCTCAAAACCTA	CTACCGGAGTCTCAAAGGGCC	ב יז (יז (ACCGGAGTCTCAAAGCGCCTG	CCGGAGTCTCAAAGCGCCTGG	CGGAGTCTCAAAGCGCCTGGG	GGAGTCTCAAAGCGCCTGGGG	GAGTCTCAAAGCGCCTGGGGC	AGTCTCAAAGCGCCTGGGGCT	GTCTCAAAGCGCCTGGGGCTC	CTCAAAGCGCC	TCAAAGCGCCTGGGGGTTCG	AAAGCGCCTGGGGCTCGC	CAAAGCGCCTGGGGCTCGCGG	AAAGCGCCTGGGGCTCGCGGA	AAGCGCCTGGGGCTCGCGGAC	AGCGCCTGGGGCTCGCGACC		֓֞֝֞֝֞֞֞֝֞֝֞֞֝֞֞֝֞֓֞֝֓֓֓֞֝֞֓֓֓֞֝֓֓֓֓֞֝֓֓֓֞֝֓֓֓֞֝֓֓֓֝֓֡֝֓֡֝֓֡ ֓֓֞֞֞֞֞֞֞֞֞֞	いしてなられていることではいいにしていることでは、	CTGGGGCTCGCGGACCCG	
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	. 0	
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	
0	0	0	0	0	0	0	Ö	0	0	0	0	0	0	0	0	0	0	0	0	0	
21	21	21	21	21	21	21		21	21	21	21	21	21	21		21	21	21	21	21	
137	138	139	140	141	142	143		145	146	147	148	149	150	151	152	53	154	55	156	.57	

FIG. 24A (9)

CTGGGGCTCGCGGACCCGGCC	TGGGGCTCGCGGACCCGGCCC	GGGGCTCGCGGACCCGGCCCA	GGGCTCGCGGACCCGGCCCAG	GGCTCGCGGACCCGGCCCAGA	GCTCGCGGACCCGGCCCAGAG	CTCGCGGACCCGGCCCAGAGG	TCGCGGACCCGGCCCAGAGGG	CGCGGACCCGGCCCAGAGGGC	GCGGACCCGGCCCAGAGGGCG	CGGACCCGGCCCAGAGGGCGG	GGACCCGGCCCAGAGGGCGGC	GACCCGGCCCAGAGGGCGGCG	ACCCGGCCCAGAGGGCGGCGG	CCCGGCCCAGAGGGCGGCGGT	CCGGCCCAGAGGGCGGCGGTG	CGGCCCAGAGGGCGGCGGTGG	GGCCCAGAGGGCGGCGGTGGC	GCCCAGAGGGCGGCGGTGGCG	CCCAGAGGGCGGCGGTGGCGG	CCAGAGGGCGGCGGTGGCGGC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0		0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
28	59	09	61	62	63	64	65	99	6 4	68	69	70	71	72	73	74	75	97	11	78

FIG. 24A (10)

						Ť														
CAGAGGGGGGGGGGCA	AGAGGCGGCGGTGGCGGCAG	GAGGGCGGCGGCAGC	AGGGCGGCGGTGGCGGCAGCT	GGGCGGCGGTGGCGGCAGCTA	GGCGGCGGTGGCGGCAGCTAC	GCGCCGCTGCCGCCAGCTACT	CGGCGGTGGCGGCAGCTACTT	GGCGGTGGCGGCAGCTACTTT	GCGGTGGCCGCCAGCTACTTTT	$\Gamma T T$	GGTGGCGGCAGCTACTTTTCT	GTGGCGGCAGCTACTTTTCTG	TGGCGGCAGCTACTTTTCTGG	GGCGGCAGCTACTTTTCTGGT	GCGCCAGCTACTTTTTCTGGTC	CGGCAGCTACTTTTCTGGTCA	GGCAGCTACTTTTCTGGTCAG	GCAGCTACTTTTCTGGTCAGG	CAGCTACTTTTCTGGTCAGGG	AGCTACTTTCTGGTCAGGGC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21				21		21		21		21	21	21	21
σ	0	<u>-</u> -	~	ന	4	Ŋ	9	7	ω	9	0	٦	~	ന	4	Ŋ	9		ω	<u>σ</u>
17	18	18	18	18	18	18	18	18	18	18	19	19	19	19	19	19	19	19	19	19

G. 24A (11)

GCTACTTTCTGGTCAGGGCT	CTACTTTCTGGTCAGGGCTC	TGGTCAGGGCTC	ACTTTTCTGGTCAGGGCTCGG	CTTTTCTGGTCAGGGCTCGGA	TTTTCTGGTCAGGGCTCGGAC	TTTCTGGTCAGGGCTCGGACA	TTCTGGTCAGGGCTCGGACAC	TCTGGTCAGGGCTCGGACACC	CTGGTCAGGGCTCGGACACCG	TGGTCAGGGCTCGGACACCGG	GGTCAGGGCTCGGACACCGGC	GTCAGGGCTCGGACACCGGCG	TCAGGGCTCGGACACCGGCGC	CAGGGCTCGGACACCGGCGCG	AGGGCTCGGACACCGGCGCGT	GGGCTCGGACACCGGCGCGTC	GGCTCGGACACCGGCGCGTCT	GCTCGGACACCGGCGCGTCTC	CTC	rct
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0.	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21			21			21			21						21		21	21
200	201	202	203	204	205	206	207	208	209	$\overline{\Box}$	211	Н	213	214	215	216	217	218	219	220

FIG. 24A (12)

CGGACACCGGCGCGTCTCTCA	GGACACCGGCGCGTCTCTCAA	GACACCGGCGCGTCTCTCAAG	ACACCGGCGCGTCTCTCAAGC	CACCGCCGCGTCTCTCAAGCT	ACCGGCGCGTCTCTCAAGCTC	CCGCCCCTCTCAAGCTCG	CGGCGCGTCTCAAGCTCGC	GGCGCGTCTCAAGCTCGCC	GCGCGTCTCTCAAGCTCGCCT	CGCGTCTCTCAAGCTCGCCTC	GCGTCTCTCAAGCTCGCCTCT	CGTCTCTCAAGCTCGCCTCTT	GTCTCTCAAGCTCGCCTCTTC	TCTCTCAAGCTCGCCTCTTCG	CTCTCAAGCTCGCCTCTTCGG	TCTCAAGCTCGCCTCTTCGGA	CTCAAGCTCGCCTCTTCGGAG	TCAAGCTCGCCTCTTCGGAGC	CAAGCTCGCCTCTTCGGAGCT	AAGCTCGCCTCTTCGGAGCTG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21			21		21		21		21		21		21	21
221	222	223	224	225	226	227	228	229	230	231	232	233	234	235	236	237	238	239	240	241

FIG. 24A (13)

							89	1/15	56											
AGCTCGCCTCTTCGGAGCTGG	GCTCGCCTCTTCGGAGCTGGA		PTCG	CGCCTCTTCGGAGCTGGAACG	PTCGGAGCTGGAACG	CCTCTTCGGAGCTGGAACGCC	CTCTTCGGAGCTGGAACGCCT	TCTTCGGAGCTGGAACGCCTG	CTTCGGAGCTGGAACGCCTGA	TTCGGAGCTGGAACGCCTGAT	AGC.	CGGAGCTGGAACGCCTGATTG	GGAGCTGGAACGCCTGATTGT	GAGCTGGAACGCCTGATTGTC		GCTGGAACGCCTGATTGTCC	GCCTGATTGTC) (SATTGTCCCC	ATTGTCCCCA
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
4	4	4	4	4	4	4	4	S	S	S	S	S	255	വ	വ	S	S	9	9	9

FIG. 24A (14)

							,													
AACGCCTGATTGTCCCCAACA	ACGCCTGATTGTCCCCAACAG	CGCCTGATTGTCCCCAACAGC	GCCTGATTGTCCCCAACAGCA	CCTGATTGTCCCCAACAGCAA	CTGATTGTCCCCAACAGCAAC	TGATTGTCCCCAACAGCAACG	GATTGTCCCCAACAGCAACGG	ATTGTCCCCAACAGCAACGGC	TTGTCCCCAACAGCAACGGCG	TGTCCCCAACAGCAACGGCGT	GTCCCCAACAGCAACGGCGTG	TCCCCAACAGCAACGGCGTGA	CCCCAACAGCAACGGCGTGAT	CCCAACAGCAACGGCGTGATC	CCAACAGCAACGGCGTGATCA	CAACAGCAACGGCGTGATCAC	AACAGCAACGGCGTGATCACG	ACAGCAACGGCGTGATCACGA	CAGCAACGGCGTGATCACGAC	AGCAACGGCGTGATCACGACG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21		21		21	21	21							21			21	21	21	21
33	24	23	9	2.2	8	9	0	7	2	3	4	Ŋ	9		ω	δ	0		2	33
_	26	26	26		26	26	27	27	27	27	27	27	27	-	27	27	28	28	28	28

FIG. 24A (15)

GCAACGGCGTGATCACGACGA	CAACGGCGTGATCACGACGAC	AACGCCTGATCACGACGACG	ACGCCTGATCACGACGACGC	CGGCGTGATCACGACGACGCC	GGCGTGATCACGACGACGCCT	GCGTGATCACGACGACGCCTA	CGTGATCACGACGCCTAC	GTGATCACGACGCCTACA	TGATCACGACGCCTACAC	GATCACGACGACGCCTACACC		ACCC	CACGACGCCTACACCCCC	ACGACGACTACACCCCCG	CGACGACGCCTACACCCCCGG	GACGACGCCTACACCCCCGGG	AC	ACC	ACCCCGGGAC	ACC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0		0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21				21		21		21		21	21	21		21	21
4	Ŋ	9	7	ω	თ	0	_	2	က	4	Ŋ	9	7	ω	<u>ი</u>	0	٦	2	ന	4
28	28	28	28	28	28		29	29	29	29	29	29	29	59	_	30	30	30	30	30

FIG. 24A (16)

CGCCTACACCCCCGGGACAGT	GCCTACACCCCCGGGACAGTA	CCTACACCCCGGGACAGTAC	CTACACCCCGGGACAGTACT	TACACCCCGGGACAGTACTT	ACACCCCGGGACAGTACTTT	CACCCCGGGACAGTACTTTT	ACCCCCGGGACAGTACTTTTA	CCCCCGGGACAGTACTTTAC	CCCCGGGACAGTACTTTACC	CCCGGGACAGTACTTTACCC	CCGGGACAGTACTTTACCCC	CGGGACAGTACTTTTACCCCC	GGGACAGTACTTTTACCCCCG	GGACAGTACTTTTACCCCCGC	GACAGTACTTTTACCCCCGCG	ACAGTACTTTTACCCCCGCGG	CAGTACTTTTACCCCCGCGGG	AGTACTTTTACCCCCGCGGGG	GTACTTTTACCCCCGCGGGG	TACTTTTACCCCCGCGGGGT
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
305	306	307	308	309	310	311	312	313	314	315	316	317	318	319	320	321	322	323	324	325

FIG. 24A (17)

ACTITIACCCCCCCGCGGGGTG	CTTTTACCCCCCCGCGGGGTGG	TTTTACCCCCCCGCGGGGTGGC	TTTACCCCCCCGCGGGGTGGCA	TTACCCCCCCGCGGGGTGGCAG	TACCCCCCCGCGGGGTGGCAGC	ACCCCCGCGGGGGTGGCAGCG	CCCCCGCGGGGGTGGCAGCGG	CCCCGCGGGGGTGGCAGCGGT	CCCGCGGGGGTGGCAGCGGTG	CCGCGGGGGTGGCAGCGGTGG	CGCGGGGTGGCAGCGGTGGA	GCGGGGTGGCAGCGGTGGAG	CGGGGGTGGCAGCGGTGGAGG	GGGGTGGCAGCGGTGGAGGT	GGGGTGGCAGCGGTGGAGGTG	GGGTGGCAGCGGTGGAGGTGC	GGTGGCAGCGGTGGAGGTGCA	GTGGCAGCGGTGGAGGTGCAG	TGGCAGCGGTGGAGGTGCAGG	AGG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21		21		21		21	21	21	21	21	21	21	21	21	21	21	21	21	21
		328	329		331		333				337			340	341	342	343	344	345	346

FIG. 24A (18)

GCAGCGGTGGAGGTGCAGGGG	CAGCGGTGGAGGTGCAGGGGG	AGCGGTGGAGGTGCAGGGGG	GCGGTGGAGGTGCAGGGGGCG	CGGTGGAGGTGCAGGGGGGG	GGTGGAGGTGCAGGGGGCGCA	GTGGAGGTGCAGGGGGCGCAG	TGGAGGTGCAGGGGCGCAGG	GGAGGTGCAGGGGCGCAGGG	GAGGTGCAGGGGGCGCAGGGG	AGGTGCAGGGGGCGCAGGGGG	GGTGCAGGGGGGCGCAGGGGG	GTGCAGGGGGCGCAGGGGGC	TGCAGGGGGCGCAGGGGGGGG	GCAGGGGGGCGCGC	CAGGGGGCGCAGGGGGGGGG	AGGGGGCGCAGGGGGGGGGGGGGG) F-	י לי לי	GGGCGCAGGGGGGGGGGGGGGGG	GGCGCAGGGGGGGCGTCACC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21		21	21	21	21	21	21	21	21	21	21	21	21	21	21
347	348	349	350	351	352		354	352	356	357	358	359	360	361	362	363	364	365	366	367

FIG. 24A (19)

GCGCAGGGGGGGGGTCACCG	CGCAGGGGGGGCGTCACCGA	GCAGGGGGGGCGTCACCGAG	CAGGGGGGGGTCACCGAGG	AGGGGGGGGTCACCGAGGA	GGGGGGGGTCACCGAGGAG	GGGGGGGTCACCGAGGAGC	GGGCGCGTCACCGAGGAGCA	GGCGCGTCACCGAGGAGCAG	GCGCCTCACCGAGGAGCAGG	CGGCGTCACCGAGGAGCAGGA	GGCGTCACCGAGGAGCAGGAG	GCGTCACCGAGGAGCAGGAGG	CGTCACCGAGGAGCAGGAGGG	GTCACCGAGGAGCAGGGC	TCACCGAGGAGCAGGGCT	CACCGAGGAGCAGGGGCTT	ACCGAGGAGCAGGGCTTC	CCGAGGAGCAGGGGCTTCG	CGAGGAGCAGGGGCTTCGC	
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0		0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21		21	21	21	21	21	21	21	21
368	369	370	371	372	373	374	375	376	377	378	379	380	381	382	383	384	382	386	387	388

FIG. 24A (20)

	,	((•	,		
ς αυ	77	0	0	0	0	0	AGGAGCAGGAGGGCTTCGCCG
390	21	0	0	0	0	0	GGAGCAGGAGGGCTTCGCCGA
391	21	0	0	0		0	GAGCAGGAGGGCTTCGCCGAC
392	21	0	0	0	0	0	AGCAGGAGGGCTTCGCCGACG
393	21	0	0	0	0	0	C
394	21	0	0	0	0	0	CAGGAGGGCTTCGCCGACGGC
395	21	0	0	0	0	0	AGGAGGCTTCGCCGACGGCT
396	21	0	0	0	0	0	GGAGGGCTTCGCCGACGGCTT
397	21	0	0	0	0	0	GAGGGCTTCGCCGACGGCTTT
398	21	0	0	0	0	0	AGGGCTTCGCCGACGGCTTTG
399	21	0	0	0	0	0	GGGCTTCGCCGACGGCTTTGT
400	21	0	0	0	0	0	GGCTTCGCCGACGGCTTTGTC
401	21	0	0	0	0	0	GCTTCGCCGACGGCTTTGTCA
402	21	0	0	0	0	0	GCTTTGTC
403	21	0	0	0	0	0	TTCGCCGACGCCTTTGTCAAA
404	21	0	0	0	0	0	TCGCCGACGGCTTTGTCAAAG
405	21	0	0	0	0	0	CGCCGACGGCTTTGTCAAAGC
406	21	0	0	0	0	0	GCCGACGGCTTTGTCAAAGCC
407	21	0	0	0	0	0	1 CA
408	21	0	0	0	0	0	CAAAGCC
409	21	0	0	0	0	0	CAAAGCCC

FIG. 24A (21)

									9	7/1	56									
ACGGCTTTGTCAAAGCCCTGG	CGGCTTTGTCAAAGCCCTGGA	GGCTTTGTCAAAGCCCTGGAC	GCTTTGTCAAAGCCCTGGACG	CTTTGTCAAAGCCCTGGACGA	TTTGTCAAAGCCCTGGACGAT	TTGTCAAAGCCCTGGACGATC	TGTCAAAGCCCTGGACGATCT	GTCAAAGCCCTGGACGATCTG	TCAAAGCCCTGGACGATCTGC	CAAAGCCCTGGACGATCTGCA	AAAGCCCTGGACGATCTGCAC	AAGCCCTGGACGATCTGCACA	AGCCCTGGACGATCTGCACAA	GCCCTGGACGATCTGCACAAG	CCCTGGACGATCTGCACAAGA	CCTGGACGATCTGCACAAGAT	CTGGACGATCTGCACAAGATG	TGGACGATCTGCACAAGATGA	GGACGATCTGCACAAGATGAA	GACGATCTGCACAAGATGAAC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
\vdash	\vdash	\vdash	Н	٦	٦	416	\vdash	ᠬ	Н	2	2	2	2	2	2	\sim	2	2	2	430

FIG. 24A (22)

							98	/15	6											
ACGATCTGCACAAGATGAACC	CGATCTGCACAAGATGAACCA	GATCTGCACAAGATGAACCAC	ATCTGCACAAGATGAACCACG	TCTGCACAAGATGAACCACGT	CTGCACAAGATGAACCACGTG	TGCACAAGATGAACCACGTGA	GCACAAGATGAACCACGTGAC	CACAAGATGAACCACGTGACA	ACAAGATGAACCACGTGACAC	CAAGATGAACCACGTGACACC	AAGATGAACCACGTGACACCC	AGATGAACCACGTGACACCCC	GATGAACCACGTGACACCCCC	ATGAACCACGTGACACCCCCC	TGAACCACGTGACACCCCCCA	GAACCACGTGACACCCCCCAA	AACCACGTGACACCCCCCAAC	ACCACGTGACACCCCCCAACG	CCACGTGACACCCCCCAACGT	CACGTGACACCCCCCAACGTG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
											21									
3	m	\sim	m	\sim	$^{\circ}$	m	m	m	4	4	442	4	4	4	4	4	4	4	S	\mathbf{D}

FIG. 24A (23)

ACGTGACACOCOCASACGTGA	では、このでは、このでは、このでは、このでは、このでは、このでは、このでは、この	りょういんけいいいいいじょう	「りょうしてないこうこうでしている」では、これでは、これでは、これでは、これでは、これでは、これでは、これでは、これ	ACACCCCCAACGIGIC	ACACCCCCAA	ACACCCCCAACGTGTCCCTG	CACCCCCAACGTGTCCCTCC		かていこう すかい さずい ファファン・ファン・ファン・ファン・ファン・ファン・ファン・ファン・ファン・ファ	プラブンシン でんじん マンフラン・ファン・ファン・ファン・ファン・ファン・ファン・ファン・ファン・ファン・ファ	つつてもてもとしゃくしつつ).T9.T9).WY	CCCAACGTGTCCCTGGGCGCT	CCAACGTGTCCCTGGGCGTA	#\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	「プランラララコンシュ	COTO TO)	CGTGTCCCTGGGCGCTACCGG		りつつは こうりつりつう こうしょうしゅうしゅうしゅうしゅうしゅうしゅうしゅうしゅうしゅう	HOODE ACCUMENT OF THE		TCCCTGGGGGCTACCGGGGGG
0	0	· C	· C	0	> (0	0	0	0) C) C) (0	0	0	· C) c	> (0	0	· C	· c	>	0
0	0	0	· C	o c) (0	0	0	0	0) C	> (0	0		C) C	> (>	0	c) C	>	0
0	0	0	C) C	> 0	>	0	0	0	0	· C	0 (>	0	0	0	C	O	>	0	C	· c) (0
0	0	0	0	· c	0 0	>	0	0	0	0	0	• •	>	0	0	0	C	· c	>	0	0	C		>
0	0	0	0	C) c	>	0	0	0	0	0	• •	> •	0	0	0	0		>	0	0	C) (>
21	21	21	21	2.1	ור	7 0	77	21	21	21	21	7 7	T 0			21	21			21		21		
	453		455		-) L	Ω	459	9	461	462	462	Ó	Ó	465		167	ανη) 1	169	170	171	77	

FIG. 24A (24)

0 cccTGGGCGCTACCGGGGGGC	0 CCTGGGCGCTACCGGGGGGCC	0 CTGGGCGCTACCGGGGGGCCC	0 TGGGCGCTACCGGGGGGCCCC	0 GGGCGCTACCGGGGGGCCCCC	0 GGCGCTACCGGGGGGCCCCCG	0 GCGCTACCGGGGGGCCCCCGG	CGCTACCGGGGG	0 GCTACCGGGGGGCCCCCGGCT	0 CTACCGGGGGGCCCCCGGCTG	0 TACCGGGGGGCCCCCGGCTGG	0 ACCGGGGGGCCCCCGGCTGGG	O CCGGGGGGCCCCCGGCTGGGC	0 ceeeeeccccceecreecc	0 GGGGGCCCCCGGCTGGGCCC	0 GGGGGCCCCGGCTGGGGCCCG	0 GGGGCCCCGGCTGGGCCCGG	0 GGCCCCCGGCTGGGCCCGGG	0 GGCCCCGGCTGGGCCCGGGG	0 GCCCCCGGCIGGGCCCCGGGGG	O CCCCCGCTGGGCCCCGGGGC
		0	0		0	0	0	0	0	0	0	0	0	0	0					0 0
				0 0																0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21		21				21			21			21	21	21	21	21	21	21
173	174	175		-	178	-	180	181	182		184	185	∞	187	488	489	490	491	492	493

FIG. 24A (25)

SCCCGGCTGGGCCCGGGGGCG	CCCGGCTGGGCCCGGGGGGCGT	CCGGCTGGGCCCGGGGGCGTC	CGGCTGGGCCCGGGGGCGTCT	GGCTGGGCCCGGGGGGCGTCTA	GCTGGGCCCGGGGGCGTCTAC	CIGGGCCCGGGGCGTCTACG	TGGCCCGGGGGCGTCTACGC	GGGCCCGGGGGCGTCTACGCC	GGCCCGGGGCGTCTACGCCG	GCCCGGGGGGTCTACGCCGG	CCCGGGGGGGTCTACGCCGGC	CCGGGGGCGTCTACGCCGGCC	CGGGGGCGTCTACGCCGGCCC	GGGGCGTCTACGCCGGCCCG	GGGCGTCTACGCCGGCCCGG	GGGCGTCTACGCCGGCCCGGA	GGCGTCTACGCCGGCCCGGAG	GCGTCTACGCCGGCCCGGAGC	CGTCTACGCCGGCCCGGAGCC	GTCTACGCCGGCCCGGAGCCA
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21				21		21					21			21		21	21	21	21	21
494	495	496	497	498	499	500	501	502	503	504		506	507	508	_	510	511	512	513	514

FIG. 24A (26)

00	0 0	0 0	0 0	0 0	
	0	o c)) C	TACGCCGGCCCACC TACGCCACC
	0	0	0	0	GCCGGCCGGAGCCA
	0	0	0	0	CCGGAGCCACCTC
	0	0	0	0	ACCTCC
	0	0	0	0	ACCTCCC
	0	0	0	0	ACCTCCCG
	0	0	0	0	GGCCCGGAGCCACCTCCCGTT
	0	0	0	0	GCCCGGAGCCACCTCCCGTTT
	0	0	0	0	CCTCCCGT
	0	0	0	0	$C_{\mathcal{O}}$
	0	0	0	0	TTTAC
	0	0	0	0	GGAGCCACCTCCCGTTTACAC
	0	0	0	0	GAGCCACCTCCCGTTTACACC
	0	0	0	0	AGCCACCTCCCGTTTACACCA
	0	0	0	0	GCCACCTCCCGTTTACACCAA
	0	0	0	0	CCGTTTACACCA
	0	0	0	0	TTACACCAAC
	0	0	0	0	TACACCAACC
	0	0	0	0	TTACACCAACC

FIG. 24A (27)

CTCCCGTTTACACCAACCTCA	TCCCGTTTACACCAACCTCAG	CCCGTTTACACCAACCTCAGC	CCGTTTACACCAACCTCAGCA	CGTTTACACCAACCTCAGCAG	AG	TTTACACCAACCTCAGCAGCT	TTACACCAACCTCAGCAGCTA	TACACCAACCTCAGCAGCTAC	ACACCAACCTCAGCAGCTACT	CACCAACCTCAGCAGCTACTC	H	C	ICC	AACCTCAGCAGCTACTCCCCA	ACCTCAGCAGCTACTCCCCAG	CCTCAGCAGCTACTCCCCAGC	CTCAGCAGCTACTCCCCAGCC	TCCCCAGC	CAGCAGCTACTCCCCAGCCTC	CCCAGCCTC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	O	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21		21				21					21						21	21	21	21
536	537	538	539	540	541	542	543	544	545	546	547	548	549	550	551	552	553	554	552	556

FIG. 24A (28)

						,														
GCAGCTACTCCCCAGCCTCTG	CAGCTACTCCCCAGCCTCTGC	AGCTACTCCCCAGCCTCTGCG	GCTACTCCCCAGCCTCTGCGT	CTACTCCCAGCCTCTGCGTC	GT	GT	CTCCCCAGCCTCTGCGTCCTC	TCCCCAGCCTCTGCGTCCTCG	CCIC	()	CCAGCCTCTGCGTCCTCGGGA	CAGCCTCTGCGTCCTCGGGAG	AGCCTCTGCGTCCTCGGGAGG	GCCTCTGCGTCCTCGGGAGGC	CCTCTGCGTCCTCGGGAGGCG	CTCTGCGTCCTCGGGAGGCGC	TCTGCGTCCTCGGGAGGCGCC	CTGCGTCCTCGGGAGGCGCCG	TGCGTCCTCGGGAGGCGCCGG	GCGTCCTCGGGAGGCGCCGGG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	O	0	0	0	0	0	0	0	0	0	.0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21		21				21													21	21
557		559	260	561		563					9	Ö	~		_	1	574	<u></u>	576	•



FIG. 24A (29)

CGTCCTCGGGAGGCGCCGGGG	GTCCTCGGGAGGCGCCGGGGC	TCCTCGGGGGCT	CCTCGGGAGGCGCCGGGGCTG	CTCGGGAGGCGCCGGGGCTGC	TCGGGAGGCGCCGGGGCTGCC	CGGGAGGCGCCGGGGCTGCCG	GGGAGGCCCGGGGCTGCCGT	GGAGGCCCGGGGCTGCCGTC	GAGGCGCCGGGCTGCCGTCG	AGGCGCCGGGCTGCCGTCGG	GGCGCCGGGCTGCCGTCGGG	GCGCCGGGCTGCCGTCGGGA	CGCCGGGGCTGCCGTCGGGAC	GCCGGGGCTGCCGTCGGGACC	CCGGGGCTGCCGTCGGGACCG	()	GGGGCTGCCGTCGGGACCGGG	GGGCTGCCGTCGGGACCGGGA	GGCTGCCGTCGGGACCGGGAG	GCTGCCGTCGGGACCGGGAGC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21			21	21		21	21	21		21	21	21	21	21	21	21	21
578	579	580	581	582	583		585	586	587	588	589	590	591	592	593	594	595	596	597	598

FIG. 24A (30)

CTGCCGTCGGGACCGGGAGCT	TGCCGTCGGGACCGGGAGCTC	GCCGTCGGGACCGGGAGCTCG	CCGTCGGGACCGGGAGCTCGT	CGTCGGGACCGGGAGCTCGTA	GTCGGGACCGGGAGCTCGTAC	TCGGGACCGGGAGCTCGTACC	CGGGACCGGGAGCTCGTACCC	GGGACCGGGAGCTCGTACCCG	GGACCGGGAGCTCGTACCCGA	GACCGGGAGCTCGTACCCGAC	ACCGGGAGCTCGTACCCGACG	CCGGGAGCTCGTACCCGACGA	CGGGAGCTCGTACCCGACGAC	GGGAGCTCGTACCCGACGACC	GGAGCTCGTACCCGACGACCA	GAGCTCGTACCCGACGACCAC	AGCTCGTACCCGACGACCACC	GCTCGTACCCGACGACCACCA	CTCGTACCCGACGACCACCAT	TCGTACCCGACGACCACCATC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
		0																		
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21		21				21		21		21		21		21	21	21	21	21
	009	601	602	603	604	605	909	607	809	609		611	612	613	614	615	919	617	618	619

FIG. 24A (31)

CGTACCCGACGACCATCA	CATCA	CCGACGACCACC	CCGACGACCACCATCAG	GACCACCATCAGC	CATCAGCTA	CACCATCAGCT	CAGCTACC	AGCTACC	ATCAGCTACC	CACCATCAGCTACCT	AGCTACCTCC	CATCAGCTACCTCC	CATCAGCTACCTCCCAC	AGCTACCTCCCAC	GCTACCTCCCACAC	CAGCTACCTCCAC	TCAGCTACCTCCACACACCC	CAGCTACCTCCCACAC	AGCTACCTCCCACAC	CTCCACACGCGC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0		0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	.0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
						21	21	21	21	21	21	21	21				21	21	21	21
20	21	22	23	24	25		27	28	60	30	31	32	33	34	ເດ	98	37	<u>∞</u>	9	0
		9			9	9		9			9	63	63	63	63	63	63	63	63	64

FIG. 24A (32)

641	21	0	0	0	0	0	GCTACCTCCCACACGCGCCGC
642	21	0	0	0	0	0	CTACCTCCCACACGCGCCGCC
643	21	0	0	0	0	0	TACCTCCCACACGCGCCGCCC
644		0	0	0	0	0	ACCTCCCACACGCGCCGCCCT
645		0	0	0	.0	0	CCTCCCACACGCGCCGCCCTT
646	21	0	0	0	0	0	CTCCCACACGCGCCGCCCTTC
647	21	0	0	0	0	0	TCCCACACGCGCCGCCCTTCG
648	21	0	0	0	0	0	CCCACACGCGCCGCCCTTCGC
649	21	0	0	0	0	0	CCACACGCCGCCCTTCGCC
650	21	0	0	0	0	0	CACACGCCGCCCTTCGCCG
651	21	0	0	0	0	0	ACACGCGCCCCTTCGCCGG
652	21	0	0	0	0	0	CACGCCCCCTTCGCCGGT
653	21	0	0	0	0	0	ACGCCCCCCTTCCCCCGGTG
654	.21	0	0	0	0	0	CGCGCCCCTTCGCCGGTGG
655	21	0	0	0	0	0	GCGCCCCTTCGCCGGTGGC
929		0	0	0	0	0	CGCCGCCCTTCGCCGGTGGCC
657	21	0	0	0	0	0	GCCGCCCTTCGCCGGTGGCCA
658		0	0	0	0	0	CCGCCCTTCGCCGGTGGCCAC
629	21	0	0	0	0	0	CGCCCTTCGCCGGTGGCCACC
099	21	0	0	0	0	0	GCCCTTCGCCGGTGGCCACCC
661	21	0	0	0	0	0	CCACCC

FIG. 24A (33)

	りのファイスのでは、このでは、このでは、このでは、このでは、このでは、このでは、このでは、こ) (りょりりし		CGCCGGCGCACCCGGCGCA	GCCGGCCACCCGGCGCAG	CCGCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	\mathbf{C}	GGTGGCCACCGGCGCAGCTG	GTGGCCACCCGGCGCACTACA	Tegananananananananananananananananananan	りりり「こうないのこうこう」	SOCCACIOCACIOCOCOCOCOCOCOCOCOCOCOCOCOCOCO	GCCACCGGCGCAGCTGGGCT	CCACCGGCGCAGCTGGGGTT	CACCCCGGCGCACACCCCCACACCCCCCCCCCCCCCCCC	の「こうののでしたないのののののでして」といっている。		CCCGCAGCTGGGCTTGGG	CCGGCGCAGCTGGGCTTGGGC	<u> </u>		SCAGCTGGGC
0	0	· C	· C	o c	> c	> 0	> 0	> 0	>	0	0	C	0	>	0	0	C	· c	>	0	0	0	0
0	0	0	· C	o c	o c	o c	O C	O (o	0	0	C) c	>	0	0	0	· c	> '	0	0	0	0
0	0	0	C	o c) C	o c) C	> C	> (0	0	0	· C	> (0	0	0	· C) (0		0	0
0	0	0	0	· C) C	o c) C	o c	> (0	0	0	· C	> 0	0	0	0	C	> 0	>	0	0	0
0	0	0	0	C) C	· C	o c	· C	> (0	0	0	C	O	>	0	0	C	• •	>	0	0	0
21	21	21		21				1 6	1 6		21	21	1			21		21		-	21	21	21
25	33	54	5	9	7	- ∞	-) _[2	സ	7		Ω ·	9	7	ω			. ·	. , H	7
99	99	99	99	99	99				•			67	67			67	67	67	-		χ	89	89

FIG. 24A (34)

683	21	0	0	0	0	0	CGCAGCTGGGCTTGGGCCGCG
684	21	0	0	0	0	0	GCAGCTGGGCTTGGGCCGCGG
685		0	0	0	0	0	CAGCTGGGCTTGGGCCGCGGC
989	21	0	0	0	0	0	AGCTGGGCTTGGGCCGCGGCG
687		0	0	0	0	0	GCTGGGCTTGGGCCGCGCGC
688		0	0	0	0	0	CTGGGCTTGGGCCGCGCGCC
689	21	0	0	0	0	0	TGGGCTTGGGCCGCGCGCCT
069		0	0	0	0	0	GGGCTTGGGCCGCGCGCCTC
691		0	0	0	0	0	GGCTTGGGCCGCGCGCCTCC
692		0	0	0	0	0	GCTTGGGCCGCGCCCTCCA
693	21	0	0	0	0	0	CTTGGGCGCGCGCCTCCAC
694		0	0	0	0	0	TIGGCCGCGCCCCTCCACC
695	21	0	0	0	0	0	TGGGCCGCGCCTCCACCT
969	21	0	0	0	0	0	GGGCCGCGCCCTCCACCTT
697	21	0	0	0	0	0	GGCCGCCGCCTCCACCTTC
869	21	0	0	0	0	0	GCCGCGCCCTCCACCTTCA
669	21	0	0	0	0	0	CCGCGCCCTCCACCTTCAA
700	21	0	0	0	0	0	CGCGGCGCCTCCACCTTCAAG
701	21	0	0	0	0	0	GCGGCGCCTCCACCTTCAAGG
702	21	0	0	0	0	0	CGGCGCCTCCACCTTCAAGGA
703	21	0	0	0	0	0	GGCGCCTCCACCTTCAAGGAG

FIG. 24A (35)

GCGCCTCCACCTTCAAGGAGG	CGCCTCCACCTTCAAGGAGGA	GCCTCCACCTTCAAGGAGGAA	CCTCCACCTTCAAGGAGGAAC	CTCCACCTTCAAGGAGGAACC	TCCACCTTCAAGGAGGAACCG	CCACCTTCAAGGAGGAACCGC	CACCTTCAAGGAGGAACCGCA	ACCTTCAAGGAGGAACCGCAG	CCTTCAAGGAGGAACCGCAGA	CTTCAAGGAGGAACCGCAGAC	TTCAAGGAGGAACCGCAGACC	TCAAGGAGGAACCGCAGACCG	CAAGGAGGAACCGCAGACCGT	AAGGAGGAACCGCAGACCGTG	AGGAGGAACCGCAGACCGTGC	GGAGGAACCGCAGACCGTGCC	GAGGAACCGCAGACCGTGCCG	AGGAACCGCAGACCGTGCCGG	GGAACCGCAGACCGTGCCGGA	GAACCGCAGACCGTGCCGGAG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	· 0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21		21		21	21	21		21		21		21		21	2.1	21	21	21	21
704	705	706	707	708	709	710	711	712	713	714	715	716	717	718	719	720	721	722	723	724

FIG. 24A (36)

AACCGCAGACCGTGCCGGAGG	Ü	CCGCAGACCGTGCCGGAGGCG	CGCAGACCGTGCCGGAGGCGC	GCAGACCGTGCCGGAGGCGCG	CAGACCGTGCCGGAGGCGCGC	AGACCGTGCCGGAGGCGCGCA	GACCGTGCCGGAGGCGCGCAG	ACCGTGCCGGAGGCGCGCAGC	CCGTGCCGGAGGCGCGCAGCC	CGTGCCGGAGGCGCGCAGCCG	GTGCCGGAGGCGCGCAGCCGG	TGCCGGAGGCGCGCAGCCGGG	GCCGGAGGCGCGCAGCCGGGA	CCGGAGGCGCGCAGCCGGGAC	CGGAGGCGCGCAGCCGGGACG	GGAGGCGCGCAGCCGGGACGC	GAGGCGCGCAGCCGGGACGCC	AGGCGCGCAGCCGGGACGCCA	GGCGCGCAGCCGGGACGCCAC	GCGCGCAGCCGGGACGCCACG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0		0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	Ö	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
25	26	27	28	59	30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45

FIG. 24A (37)

CGCGCAGCCGGGACGCCACGC	GCGCAGCCGGGACGCCACGCC	CGCAGCCGGGACGCCACGCCG	GCAGCCGGGACGCCACGCCGC	CAGCCGGGACGCCACGCCGCC	AGCCGGGACGCCACGCCGCCG	GCCGGGACGCCACGCCGCCGG	CCGGGACGCCACGCCGCCGGT	CGGGACGCCACGCCGCCGGTG	GGGACGCCACGCCGCCGGTGT	GGACGCCACGCCGGGTGTC	GACGCCACGCCGCCGGTGTCC	ACGCCACGCCGCCGGTGTCCC	CGCCACGCCGCCGGTGTCCCC	GCCACGCCGCTGTCCCCC	CCACGCCGCTGTCCCCCA	CACGCCGCCGGTGTCCCCCAT	ACGCCGCCGGTGTCCCCCATC	CGCCGCCGGTGTCCCCCATCA	GCCGCCGGTGTCCCCCATCAA	CCGCCGGTGTCCCCCATCAAC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21		21	21	21		21	21	21		21	21	21	21	21
746	747	748	749	750	751	752	753	754	755	756	757	758		160	761	762	763	764	765	992

(38)	CGCCGGTGTCCCCCATCAACA	CCCCATCAAC	CCGGTGTCCCCCATCAACATG	CGGTGTCCCCCATCAACATGG	GGTGTCCCCCATCAACATGGA	GTGTCCCCCATCAACATGGAA	TGTCCCCCATCAACATGGAAG	GTCCCCCATCAACATGGAAGA	TCCCCCATCAACATGGAAGAC	CCCCCATCAACATGGAAGACC	CCCCATCAACATGGAAGACCA	CCCATCAACATGGAAGACCAA	CCATCAACATGGAAGACCAAG	CATCAACATGGAAGACCAAGA	ATCAACATGGAAGACCAAGAG	TCAACATGGAAGACCAAGAGC	CAACATGGAAGACCAAGAGCG	AACATGGAAGACCAAGAGCGC	ACATGGAAGACCAAGAGCGCA	CATGGAAGACCAAGAGCGCAT	ATGGAAGACCAAGAGCGCATC
FIG. 24A (38)	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
FIG.	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
		21																21	21	21	21
		9	169	_		7	773	774	775	9//	777	778	779	780	∞	∞		ω	785	982	787

FIG. 24A (39)

TGGAAGACCAAGAGCGCATCA	GGAAGACCAAGAGCGCATCAA	GAAGACCAAGAGCGCATCAAA	AAGACCAAGAGCGCATCAAAG	AGACCAAGAGCGCATCAAAGT	GACCAAGAGCGCATCAAAGTG	ACCAAGAGCGCATCAAAGTGG	CCAAGAGCGCATCAAAGTGGA	CAAGAGCGCATCAAAGTGGAG	AAGAGCGCATCAAAGTGGAGC	AGAGCGCATCAAAGTGGAGCG	GAGCGCATCAAAGTGGAGCGC	AGCGCATCAAAGTGGAGCGCA	GCGCATCAAAGTGGAGCGCAA	CGCATCAAAGTGGAGCGCAAG	GCATCAAAGTGGAGCGCAAGC	CATCAAAGTGGAGCGCAAGCG	ATCAAAGTGGAGCGCAAGCGG	TCAAAGTGGAGCGCAAGCGGC	CAAAGTGGAGCGCAAGCGGCT	AAAGTGGAGCGCAAGCGGCTG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0		0	0	0	0	0	0	0	0	0	0	O	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
788	789	190	791	792	793	794	795	962	797	798	799	800	801	802	803	804	802	908	807	808

FIG. 24A (40)

AAGTGGAGCGCAAGCGGCTGC	AGIIGGAGCGCAAGCGGCIIGCG	GIGGAGCGCAAGCGGCIGGG	TGGAGCGCAAGCGGCTGCGGA	GGAGCGCAAGCGGCTGCGGAA	GAGCGCAAGCGGCTGCGGAAC	AGCGCAAGCGGCTGCGGAACC	GCGCAAGCGGCTGCGGAACCG	CGCAAGCGGCTGCGGAACCGG	GCAAGCGGCTGCGGAACCGGC	CAAGCGGCTGCGGAACCGGCT	AAGCGGCTGCGGAACCGGCTG	AGCGGCTGCGGAACCGGCTGG	GCGGCTGCGGAACCGGCTGGC	CGGCTGCGGAACCGGCTGGCG	GGCTGCGGAACCGGCTGGCGG	GCTGCGGAACCGGCTGGCGGC	CTGCGGAACCGGCTGGCGGCC	TGCGGAACCGGCTGGCGGCCA	GCGGAACCGGCTGGCGGCCAC	CGGAACCGGCTGGCGGCCACC
0.0	o (> (0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0 (၁	0 (0	0		0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	> (0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0 (0)	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	> (0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21		7.7 7.7		21	21	21	21	21	21	21	21	21	21		21	21	21	21	21	21
809	810	- 1 ,	812	813	814	815	816	817	818	819	820	821	822	823	824	825	826	827	828	829

FIG. 24A (41)

GGAACCGGCTGTTTTTTTTTTTTTTTTTTTTTTTTTTTT	GAACCGGCTGGCGGCCALOAD	AACCGGCTGGCGCCACCAAG	ACCGGCTGGCGGCGACA			GGCTGGCGGCCACCAAGTGCC	GCTGGCGGCCACCAAGTGCCG	CTGGCGGCCACCAAGTGCCGG	TGGCGGCCACCAAGTGTCA	GGCGGCCACCAAGTGCCGGAA	GCGCCACCAAGTGCCCAAA			GCCACCAAGTGCCGGAAGCGG	CCACCAAGTGCCGGAAGCGGA	CACCAAGTOTOTOTOTOTOTOTOTOTOTOTOTOTOTOTOTOTOT	ACCA A G G C C C C C C C C C C C C C C C		いっては、そのことであることでは、これでは、そのことであることでは、そのことできません。	AAGTGCCGGAAGCGGAAGCT
0	0	0	0	0	.0	0	0	0	0	0	0	0	0	0	0	0	0	· C	· C	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	C) C	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
830	831	832	833	834	835	836	837	838	839	840	841	842	843	4	845	846	847	848	849	850

FIG. 24A (42)

						/	18/	156	,											
AGTGCCGGAAGCGGAAGCTGG	GTGCCGGAAGCGGAAGCTGGA	TGCCGGAAGCGGAAGCTGGAG	GCCGGAAGCGGAAGCTGGAGC	CCGGAAGCGGAAGCTGGAGCG	CGGAAGCGGAAGCTGGAGCGC	GGAAGCGGAAGCTGGAGCGCA	GAAGCGGAAGCTGGAGCGCAT	AAGCGGAAGCTGGAGCGCATC	AGCGGAAGCTGGAGCGCATCG	GCGGAAGCTGGAGCGCATCGC	CGGAAGCTGGAGCGCATCGCG	GGAAGCTGGAGCGCATCGCGC	GAAGCTGGAGCGCATCGCGCG	AAGCTGGAGCGCATCGCGCGC	AGCTGGAGCGCATCGCGCGCC	GCTGGAGCGCATCGCGCGCCT	CTGGAGCGCATCGCGCGCCTG	TGGAGCGCATCGCGCGCCTGG	GGAGCGCATCGCGCGCCTGGA	GAGCGCATCGCGCGCCTGGAG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	O	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
\mathcal{L}	S	S	Ω	S	S	Ŋ	S	S	9	9	9	9	9	9	9	9	9	9	870	7

FIG. 24A (43)

0 0 AGCGCATCGCGCCCTGGAGG	0	0	0	0	0	0	0	0	0	0 0 GCGCCTGGAGAAGGTGAA	0	0	0	0	0	0	0	0	0 0 GGACAAGGTGAAGACGCTCAA	
0 0																				•
0	0	0	0																0	C
0	0		0							0										
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	C
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	נכ
872	873	874	875	876	877	878	879	880	881	882	883	884	882	886	887	888	889	890	891	892

'G. 24A (44)

ACAAGGTGAAGACGCTCAAGG	CAAGGTGAAGACGCTCAAGGC	AAGGTGAAGACGCTCAAGGCC	AGGTGAAGACGCTCAAGGCCG	GGTGAAGACGCTCAAGGCCGA	GTGAAGACGCTCAAGGCCGAG	TGAAGACGCTCAAGGCCGAGA	GAAGACGCTCAAGGCCGAGAA	AAGACGCTCAAGGCCGAGAAC	AGACGCTCAAGGCCGAGAACG	GACGCTCAAGGCCGAGAACGC	ACGCTCAAGGCCGAGAACGCG	CGCTCAAGGCCGAGAACGCGG	GCTCAAGGCCGAGAACGCGGG	CTCAAGGCCGAGAACGCGGGG	TCAAGGCCGAGAACGCGGGGC	CAAGGCCGAGAACGCGGGGCT	AAGGCCGAGAACGCGGGGCTG	AGGCCGAGAACGCGGGGCTGT	GGCCGAGAACGCGGGGCTGTC	GCCGAGACGCGGGGCTGTCG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	O	0		0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0		0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
93 21	94 21	95 21	96 21	97 21	98 21	σ	00 21	01 21	02 21	03 21	04 21	05 21	06 21	7 2	8	09 21	10 21		12 21	13 21
ω	ω	ω	ά	ά	ώ							ŏ			-	_	<u>o</u> ,	9	<u>,</u>	9

FIG. 24A (45)

						*														
CCGAGAACGCGGGGCTGTCGA	CGAGAACGCGGGGCTGTCGAG	GAGAACGCGGGCTGTCGAGT	AGAACGCGGGCTGTCGAGTA	GAACGCGGGCTGTCGAGTAC	AACGCGGGCTGTCGAGTACC	ACGCGGGCTGTCGAGTACCG	CGCGGGCTGTCGAGTACCGC	GCGGGCTGTCGAGTACCGCC	CGGGGCTGTCGAGTACCGCCG	GGGCTGTCGAGTACCGCCGG	GGGCTGTCGAGTACCGCCGGC	GGCTGTCGAGTACCGCCGGCC	GCTGTCGAGTACCGCCGGCCT	CTGTCGAGTACCGCCGGCCTC	TGTCGAGTACCGCCGGCCTCC	GTCGAGTACCGCCGGCCTCCT	TCGAGTACCGCCGGCCTCCTC	CGAGTACCGCCGGCCTCCTCC	GAGTACCGCCGGCCTCCTCCG	AGTACCGCCGGCCTCCTCCGG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	.0	o [.]	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21		21		21	21	21	21	21	21	21	21	21	21
14	15	16	17	18	19	2.0	21	22	23	24	25	26	27	28	29	30	31	32	33	34
9	9	9	9	9	g	9	9	9	9	9	9	9	9	9	9	9	9	9	9	9

FIG. 24A (46)

					12	æ/.	156													
GTACCGCCGGCCTCCTCCGGG	CCTCCTCGGG	Ü	GGGAG	CGCCGGCCTCCTCCGGGAGCA	CA	CCGGCCTCCTCCGGGAGCAGG	CGGCCTCCTCCGGGAGCAGGT	GGCCTCCTCCGGGAGCAGGTG	GCAGGT	CCTCCTCCGGGAGCAGGTGGC	SCAGGTGGC	TGGCC	GTGGCCC	\mathcal{C}	CAG	CCGGGAGCAGGTGGCCCAGCT	\circ	GGCCCAGCTC	TGGCCCAGCTCA	CCAGCTCA
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
3	\sim	$^{\circ}$	938	\sim	4	4	4	4	4	4	4	4	4	4	വ	വ	S	D	Ω	Ω

FIG. 24A (47)

					/	23,	/15	6												
AGCAGGTGGCCCAGCTCAAAC	GCAGGTGGCCCAGCTCAAACA	CAGGTGGCCCAGCTCAAACAG	AGGTGGCCCAGCTCAAACAGA	GGTGGCCCAGCTCAAACAGAA	GTGGCCCAGCTCAAACAGAAG	TGGCCCAGCTCAAACAGAAGG	GGCCCAGCTCAAACAGAAGGT	GCCCAGCTCAAACAGAAGGTC	CCCAGCTCAAACAGAAGGTCA	CCAGCTCAAACAGAAGGTCAT	CAGCTCAAACAGAAGGTCATG	AGCTCAAACAGAAGGTCATGA	GCTCAAACAGAAGGTCATGAC	CTCAAACAGAAGGTCATGACC	TCAAACAGAAGGTCATGACCC	CAAACAGAAGGTCATGACCCA	AAACAGAAGGTCATGACCCAC	AACAGAAGGTCATGACCCACG	ACAGAAGGTCATGACCCACGT	CAGAAGGTCATGACCCACGTC
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0		0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
2	Ω	S	S	9	9	962	9	9	9	9	9	9	9	7	7	7	7	7	7	7

FIG. 24A (48)

							•													
AGAAGGTCATGACCCACGTCA	GAAGGTCATGACCCACGTCAG	AAGGTCATGACCCACGTCAGC	AGGTCATGACCCACGTCAGCA	GGTCATGACCCACGTCAGCAA	GTCATGACCCACGTCAGCAAC	TCATGACCCACGTCAGCAACG	CATGACCCACGTCAGCAACGG	ATGACCCACGTCAGCAACGGC	TGACCCACGTCAGCAACGGCT	GACCCACGTCAGCAACGGCTG	ACCCACGTCAGCAACGGCTGT	CCCACGTCAGCAACGGCTGTC	CCACGTCAGCAACGGCTGTCA	CACGTCAGCAACGGCTGTCAG	ACGTCAGCAACGGCTGTCAGC	CGTCAGCAACGGCTGTCAGCT	GTCAGCAACGGCTGTCAGCTG	TCAGCAACGGCTGTCAGCTGC	CAGCAACGGCTGTCAGCTGCT	AGCAACGGCTGTCAGCTGCTG
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	· O	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
21	21	21	21	21	21	21	21	21	21		21	21	21	21	21	21	21	21	21	21
977	978	979	980	981	982	983	984	985	986	987	988	686	066	991	992	993	994	995	966	266

FIG. 24A (49)

FIG. 24A (50)

016	21	0	0	0	0	0	TGCTTGGGGTCAAGGGACACG	
.017	21	0	0	0	0	0	GCTTGGGGTCAAGGGACACGC	
.018	21	0	0	0	0	0	CTTGGGGTCAAGGGACACGCC	
019	21	0	0	0	0	0	TTGGGGTCAAGGGACACGCCT	
.020	21	0	0	0	Ö	0	TGGGGTCAAGGGACACGCCTT	
.021	21	0	0	0	0	0	GGGGTCAAGGGACACGCCTTC	
.022	21	0	0	0	0	0	GGGTCAAGGGACACGCCTTCT	
.023	21	0	0	0	0	0	GGTCAAGGGACACGCCTTCTG	
024	21	0	0	0	0	C	GTCAAGGGACACGCCTTCTGA	

FIG. 24B (1)

pages)
190
of
pages
10
ŀ
File
(Partial

OligoProbe DesignStation	esignst	ation							
Probes: Preparation:	C:\HI'C	<pre>C:\HITACHI\HUMBJUNX.CDS C:\HITACHI\JUNMIX.PRP</pre>	X.CDS PRP						
Loci	rocus pos	Tm	Locu	Locus pos	Tm			rocus pos	
atqtqcacta	aaatqqa	atqtqcactaaaatqqaacaqcccttctac							
1 30	, L	1 1 2	2	2	2	7	က	4	
xunjamny	nx 1	60.76							
xunjasm	nx 1	50.03							
muscju	nx 1	30.07							
musdjunx 72	nx 721	27.84							

FIG. 24B (2)

4	4
т	м
0	~
7	7
0	7
۰ -	% -
tgtgcactaaaatggaacagcccttctac 2 29 1 1 2 humbjunx 65533 60.68 musbjunx 65533 49.58 muscjunx 1 29.97 musdjunx 721 27.66	gtgcactaaaatggaacagcccttctac 3 28 1 1 2 humbjunx 65533 60.60 musbjunx 65533 49.10 muscjunx 1 29.86 musdjunx 721 27.47

FIG. 24B (3)

	4			4				
	m			m				
	7			7				
	7			2				
	2			7				
	7			2				
CC	Н	Ç)	 1				
ccttcts	1 60.60 46.57 29.86	27.47	2000	٦	60.51	45.96	29.75	27.26
acago	5533 5533	729	0000	← 1				729
tgcactaaaatggaacagcccttctacc	4 28 1 humbjunx 60 musbjunx 60	π	godocadacygad	5 27 1	humbjunx 5	3 xunjqsnw	muscjunx 1	

FIG. 24B (4)

4	4	4
m	м	т
N	м	М
0	0	7
7	7	7
0	N	8
cactaaaatggaacagcccttctaccac 6 28 1 1 1 1 humbjunx 1 60.60 musbjunx 5 46.42 muscjunx 1 30.79 muscjunx 1 30.79 muscjunx 729 27.47	actaaaatggaacagcccttctaccacg 7 28 1 1 1 1 1 humbjunx 1 60.60 musbjunx 5 46.42 muscjunx 1 33.32 muscjunx 729 27.47	ctaaaatggaacagccttctaccacg 8 27 1 1 1 humbjunx 1 60.51 musbjunx 5 45.96 muscjunx 1 33.33 musdjunx 729 27.26

FIG. 24B (5)

4	4	4
м	m	м
ო	m .	т
2	0	7
0	73	8
7	73	α.
taaaatggaacagccttctaccacgac 9 28 1 1 1 2 humbjunx 9 60.60 musbjunx 5 49.10 muscjunx 9 34.39 muscjunx 729 27.47	aaaatggaacagccttctaccacgac 10 27 1 1 1 2 humbjunx 5 60.51 musbjunx 5 49.70 muscjunx 9 34.44 musdjunx 729 27.26	aaatggaacagccttctaccacgac 11 26 1 1 2 humbjunx 5 60.42 musbjunx 5 49.19 muscjunx 9 34.50 musdjunx 729 27.04
taaaatggaaca 9 28 humbjunx musbjunx muscjunx muscjunx	aaaatggaacagc 10 27 humbjunx musbjunx muscjunx muscjunx	aaatggaacagc 11 26 humbjunx musbjunx musbjunx muscjunx

(9)	
B	

4	Ψ
т	ഗ
т	4
N	~
Ν	N
Ν .	N
N	~
taccacgac 1 1 60.32 48.64 34.56 26.80	accacgac 1 1 60.20 48.04 34.62 32.46 33 30.25
aatggaacagccttctaccacgac 12 25 1 1 1 humbjunx 5 60.32 musbjunx 5 48.64 muscjunx 9 34.56 muscjunx 9 26.80	atggaacagccttctaccacgac 13 24 1 1 humbjunx 13 60.20 musbjunx 13 48.04 muscjunx 9 34.62 musdjunx 1 32.46 humdjunx 65533 30.25 musdjunx 737 26.55

FIG. 24B (7)

v	9
ហ	ហ
т	m
7	N
7	0
7	0
Н	0
acgac 1 1 60.08 47.39 33.39 31.14 28.83	cgacg 1 1 1 61.86 49.17 32.09 29.83 28.53
tctaccc 1 9 9 9 1 1 65533	ctaccac 1 9 9 9 1 1 65533
tggaacagcccttctaccacgac 14 23 1 1 humbjunx 9 60.00 musbjunx 9 47.3 muscjunx 9 33.3 musdjunx 1 31.1 humdjunx 65533 28.8 musdjunx 737 26.2	ggaacagcccttctaccacgacg 15 23 1 1 humbjunx 9 61.8 musbjunx 9 49.1 muscjunx 9 32.0 muscjunx 1 29.8 humdjunx 65533 28.5

FIG. 24B (8)

	ω									∞							
	4									9							
	7	26.27								7	26.27						
	2	26								7	26						
	2	737								2	737						
	2	musdjunx								2	musdjunx						
		musd									musd						
	Н									٦							
cga	٦	80.09	47.39	30.00	29.66	27.57	26.27	6.27	gac	٦	60.08	47.39	30.00	29.66	29.35	9.35	7.57
acga	Н	Ö	4	m		7		26	cgaci	۲	Ō	4	m	7		5	27
tacc	٦	σ	σ	σ	65533	Ч	281	281	acca	7	17	17	17	വ	281	281	Н
cttc	٣	humbjunx	musbjunx	muscjunx	numdjunx	musdjunx	humbjunx	musbjunx	ttct	3	humbjunx	musbjunx	muscjunx	junx	xun jqmnu	musbjunx	musdjunx
gaacagcccttctaccacgacga	16 23	humb	gsnw	musc	humd	musd	humb	musb	aacagcccttctaccacgacgac	17 23	humb	musb	musc	humdj	humb	musb	musd
gaac									aace								

FIG. 24B (9)

	œ									7							
	9									9							
	7	.27								7							
	7	26								7							
	7	737								7							
	7	musdjunx 737								7	•						
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acgacı	1 1	60.09	47.39	30.00	29.66	29.35	29.35	27.57	cgactc	1	61.86	49.17	30.00	29.66	29.35	29.35	27.57
Cacg	7	13	17	17	Ŋ	281	281	٦	acga	-1	13	17	17	വ	281	281	-
acadecerreraceaegaegae	18 23	humbjunx	musbjunx	muscjunx	humdjunx	humbjunx	musbjunx	musdjunx	cagccettetaccacgacgacte	19 23	humbjunx	musbjunx	muscjunx	humdjunx	humbjunx	musbjunx	musdjunx

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FIG. 24B (11)

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ctcatac	1 1	60.20	43.66	31.67	30.13	29.80	29.50	24.84	tcatacac	1 1	60.32	40.56	35.76	30.24	29.64	27.08
cccttctaccacgacgactcatac	22 24 1	humbjunx 17	musbjunx 17	humbjunx 281	muscjunx 17	humdjunx 5	musbjunx 281	musdjunx 5	ccttctaccacgacgactcatacac	23 25 1	humbjunx 17	musbjunx 17	humbjunx 289	muscjunx 17	musbjunx 289	humdjunx 5

FIG. 24B (12)

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cttctaccacgactcatacacag 24 26 1 1 1 humbjunx 17 60.42 musbjunx 17 44.00 humbjunx 289 35.65 musbjunx 289 29.77	ttctaccacgacgactcatacacagc 25 26 1 1 1 humbjunx 25 60.42 musbjunx 25 46.73 humbjunx 289 35.65 musbjunx 289 29.77	tctaccacgacgactcatacacagc 26 25 1 1 1 humbjunx 21 60.32 musbjunx 25 46.08 humbjunx 289 35.76 musbjunx 289 29.64

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acacagc 1 1 60.20 45.37 35.87 29.50	cacagctac 1 1 60.42 42.26 35.65 29.77	acagctac 1 1 60.32 42.64 35.76 29.64
ctaccacgacgactcatacacacacacacacacacacaca	taccacgacgactcatacacagct 28 26 1 1 1 humbjunx 21 60.42 musbjunx 25 42.26 humbjunx 289 35.65 musbjunx 289 29.77	accacgacgactcatacacagctac 29 25 1 1 1 humbjunx 29 60.32 musbjunx 25 42.64 humbjunx 289 35.76 musbjunx 289 29.64

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	ccacga 30 hu mu hu mu	cacgac 31 hu mu mu mu hu	acgacg 32 hu mu hu hu hu

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cgacgactcatacacagctacgg 33 23 1 1 1 1 humbjunx 33 60.08 musbjunx 33 41.82 humdjunx 573 26.27	gacgactcatacacagctacggg 34 23 1 1 1 humbjunx 29 60.08 musbjunx 29 41.82 humdjunx 581 26.27	acgactcatacacagctacgggatac 35 26 1 1 1 humbjunx 29 60.42 musbjunx 29 44.26 humdjunx 581 27.04
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	ctcatacacagctacgggatacgg 39 24 1 1 1 humbjunx 33 60.20 musbjunx 37 42.70 humdjunx 581 26.55	tcatacacagctacgggatacggc 40 24 1 1 1 humbjunx 33 60.20 musbjunx 37 39.75 humdjunx 581 26.55	catacacagctacgggatacggc 41 23 1 1 1 humbjunx 41 60.08 musbjunx 37 38.91 humdjunx 581 26.27

FIG. 24B (18)

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atacacagctacgggatacggcc 42 23 1 1 1 humbjunx 37 60.08 musbjunx 37 38.91 humdjunx 589 26.27	tacacagctacgggatacggccg 43 23 1 1 1 humbjunx 37 61.86 musbjunx 37 41.82 humdjunx 589 26.27	acacagctacgggatacggccg 44 22 1 1 1 humbjunx 37 61.81 musbjunx 37 40.86 humdjunx 589 25.96

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	0	70
	N	0
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FIG. 24B (19)	72	7
9	ਜ਼	
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	acagctacggatacggccg 45 21 1 1 1 humbjunx 45 61.76 musbjunx 45 40.00 humdjunx 589 25.62	cagctacgggatacggccgg 46 21 1 1 1 humbjunx 41 61.76 musbjunx 45 43.38 humdjunx 589 25.62

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agctacgggatacggccggg 48 20 1 1 1 humbjunx 41 61.70 musbjunx 45 40.35 muscjunx 561 31.40



FIG. 25

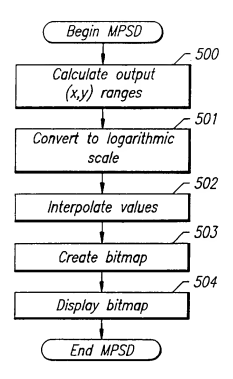
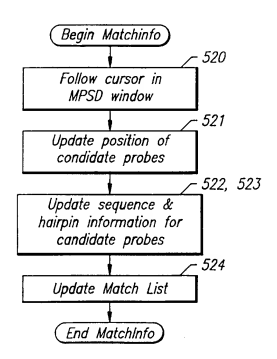


FIG. 26



	148/156	
19-DEC-1991	C TACGGGATAC G GAGCCTGGCG G ACCCGGCCCA G CGCGTCTCTC G CGCTGATCACG G CGCTGATCACG G CGCTGATCACG G CGCTGCGGCCT G CGCTGCTCCC G CGCTGCTCCC G CGCTGCTCCC G CGCTGCTCCC G CGCTGCTCCC G CGCTGCTCCCC G CGCCGGGCT G CGCCGCGCCC G CACCTTCAAG C CACCTTCCAAG C C C C C C C C C C C C C C C C C C C	
19	CATACACAGG TCCTGAAACC GGGCTCGCGG CGGACACGG ACAGCAACGG GCGTTCGCCAACGT CCCCCAACGT CCCCCAACGT CCCCCAACGT CCCCCAACGT CCCCCAACGT CCGCCCCGAACGT CCCCCAACGT CCCCCAACGT CCCCCAACGT CCCCCAACGT CCCCCAACGT CCCCCAACGT CCCCCCAACGT CCCCCCAACGT CCCCCCAACGT CCCCCCAACGT CCCCCCAACGT CCCCCCCCCC	
141 T	CACGACGACT GACTACAAAC AAAGCGCCTG GGTCAGGGCT ATTGTCCCCA TACCCCCGCG GAGCAGGAGG CACGTCTACG GCCTCTGCGT ACCACCATCA GCCTCTGCGT ACCACCATCA GCCTCTGCGC ACCACCATCA ACCACCATCA CACGTCTGCGC ACCACCATCA CACCTCTGCGC ACCACCATCA CACCTCTGCGCC ACCACCATCA CACCTCTGCGCC ACCACCATCA CACCTCTGCGCC CACCTCTGCGCC CACCTCTGCGCC CACCTCTGCGCC CACCTCTGCCCC CACCTCTGCGCC CACCTCTCGCCC CACCTCTCGCCCC CACCTCTCCCCCCCC CACCTCCCCCCCCCC	
DNA 340 G	GCCCTTCTACA CTCTCTACAC CCGGAGTCTC CTACTTTTCT GGAACGCCTG ACAGTACTTT CGTCACCGAG CTACTCCCCA GTACCCCGAG GTACCCCGAG GGCGCAGCTG GGCGCAGCTG GCGCAGCTG GCGCAGCTG GCGCATCAAA GCGGGGCTG	
1044 bp A 368 C	AAATGGAACA CTGGTGGCCT CCGACCCCTA GTGGCGGCAG CTTCGGAGCT CACCCCGGG ACGATCTGCA ACGATCTGCA ACGATCTGCA ACGATCTGCA ACGACCGGG ACCTCAGCA ACGCCCAAGA AGGCCGGAA AGGCCGGGAA AGGCCGAAGA AGGCCGAAGA AGGCCGAAAA AGGCCGAAAA AGGCCGAAAA	
HUMBJUNX 195	ATGTGCACTA GGCCGGGCCC GTCAACCTGG GAGGGCGCCCT ACGACGCCTA ACGACGCCTA GCAGGGGGCG AAAGCCCTGG GTTTACACCTGG GTTTACACCCA GCCGTCGGGA CCCTTCGCCC ATCAACATGG GAGGAACCGC ATCAACATGG GAGGCCCACCA GCGGCCCACCA GCGGCCCACCA GCGGCCCACCA GCGGCCCACCA GGGGCCCACCA GGGGCCCAGGC GGGGCCCAGGC	
US E COUNT GIN	1051 1051 1051 1051 1051 1051 1061 1061	

FIG. 28 (1)

	149/156
19-DEC-1991	TACGGGATAC GAGCCTGGCG ACCCGGCCCA CGCGTCTCTC CGTGATCACG CGCTTTTGTC GGCCTTTGTC GCCCGGGGCT ACACGCGCCC GCCCGGGGCT ACACGCGCCC GCCCGGGGCT CGCCGGGGCT CGCCGGGGCT CGCCGGGGCT CGCCGGGGCT CGCCGGGGCT CGCCGGGCCC CGCCGGGGCT CGCCGGGCCC CGCCGGGCCC CCCCCCCC
	CATACACAGC TCCTGAAACC GGGCTCGCGG ACAGCAACGG ACAGCAACGG GCTTCGCCGA CCCCCAACGT CCGGCCCGGA CCCCCAACGT CCGGCCCGCA CCCCCAACGT CCGGCCCCGA CCCCCAACGT CCGGCCCCGA CCCCCAACGT CCGGCCCCGA CCCCCCAACGT CCCCCCAACGT CCCCCCAACGT CCCCCCAACGT CCCCCCAACGT CCCCCCAACGT CCCCCCCCAACGT CCCCCCCAACGT CCCCCCCCAACGT CCCCCCCCCC
141 T	CACGACGACT GACTACAAAC AAAGCGCCTG GGTCAGGGCT ATTGTCCCCA TACCCCCGCG GAGCAGGAGG CACGTGACAC GGCTTTACG GCCTTTACG GCCTTGCGCT ACCACCATCA GCCTTGGGCC ACCACCATCA GCCTTGGGCC ACCACCATCA CGCTTGGGCC ACCACCATCA CGCTTGGGCC ACCACCATCA CGCTTGGGCC ACCACCATCA CGCCTTGGGCC ACCACCATCA CGCCTTGGGCC ACCACCATCA CGCCTTGGGCC ACCACCATCC ACCACCATCC CACGTCAGCC CACGTCAGCC
DNA 340 G	GCCCTTCTAC CTCTCTACAC CCGGAGTCTC CTACTTTTCT GGAACGCCTG ACAGTACTTT CGTCACCCAG CAAGATGAAC TGGCCCGGG CTACTCCCCA GTACCCGACG GGCGCATCAAA GCGCATCAAA GCGCATCAAA GCGCATCAAA GCGCATCAAA
1044 bp A 368 C	AAATGGAACA CTGGTGGCCT CCGACCCCTA GTGGCGGCGG CACCCCGGG CACCCCCGGG ACGTCTGCA GGCCCCCGGC ACGTCTCC ACGTCCC ACGTCCC ACCTCAGCAGA AGACCGTGCC AGACCGTGCC AGACCGTGCC AGACCGTGCC AGACCGAGA AGCCCGAGA AGCCCGAGA AGCCCAAGA AGCCCAAGA AGCCCAAGA AGCCCAAGA AGCCCAAGA AGCCCAAGA AGCCCAAGA AGCCCAAGA AGCCCAAGA
HUMBJUNX I 195	ATGTGCACTA GGCCGGGCCC GTCAACCTGG GAGGGGGCG AAGCTCGCCTA ACGACGCCTA GCAGGGGCG GTTTACACCA GCCTTCGCCG GTTTACACCA GCCGTCGGGA GCCGTCGGGG GTTTACACCA GCGGCCACCA GCGGCCACCA GCGGCCACCA GCGGCCACCA GCGGCCCACCA GCGGCCCACCA GCGGCCCACCA GCGGCCCACCA GCGGCCCACCA
LOCUS BASE COUNT ORIGIN	121 121 181 241 301 361 481 721 721 781 901

FIG. 28 (2)

19-DEC-1991	CETTCCTCCCG		_													
19	TCAACGCCTC	GCCCCAAGAA	CCGAGCTGGA	CCACCCAGTT	TCGTGCGCGC	CGCAGCCGGT	GCGGCAGCGG	GCAACTTCAA	GCCTGGCCTT	AGATGCCCGT	AGATGCCCGG	AGGCGGAGAG	TGGAGAGAAT	TGGCGTCCAC	ACCACGTTAA	
129 T	GACGATGCCC	CCGCACCTCC	CTGGCGTCGC	ACGCCGACCC	GCCGAGGGCT	ACGTCGGCGG	GCAGGGGGCA	GCAAACCTCA	9009909099	CTGCCCCAGC	ACAGTGCCCG	GAGCGGATCA	AAAAGGAAGC	AACTCGGAGC	AAAGTCATGA	TTTTGA
DNA 299 G	GACCTTCTAT	GAGCCTGAAG	GCTGCTCAAG	CATCACCACC	GGAGGGGTTC	GCCCAGCGTC	AGCCTCGGTG	GCCGGTCTAC	GCCCTCCTAC	GCCGCACCAC	GGAGCCTCAG	GGAGTCCCAG	CAAGTGCCGA	GAAAGCTCAG	GCTTAAACAG	GTTGCAAACA
996 bp A 342 C	AGATGGAAAC GACCTTATGG	ACCCAGTGGG	CCGACGTGGG	GCAACGGGCA	CAGATGAGCA	AGAACACGCT	CTCCCGCGGT	ACAGCGAGCC	ວອອອອວອອວອ	AGCAGCAGCC	CCCTGAAGGA	CCATCGACAT	TCGCTGCCTC	TGAAAACCTT	AGGTGGCACA	TAACGCAGCA
HUMCJUNX C 226	ATGACTGCAA TCCGAGAGCG	AACCTGGCCG	CTCACCTCGC	ATCCAGTCCA	AAGAACGTGA	CTGCACAGCC	GGCATGGTGG	GCCAGCCTGC	CTGAGCAGCG	CCCCAGCAGC	CGGCTGCAGG	CCCCTGTCCC	AGGAACCGCA	GAGGAAAAAG	CTCAGGGAAC	CAACTCATGC
LOCUS BASE COUNT ORIGIN	1		∞	4	0	9	\sim	∞	541	0	9	721	∞	841	901	196

FIG. 28 (3)

151/156					
24-MAY-1991		Mammalia; Hominidae.			
PRI 2		Vertebrata; Catarrhini;	·		
HUMDJUNX 1044 bp ss-mRNA Human junD mRNA	Abbeal jun-D gene; oncogene. Homo sapiens RNA.	w •• ⊃	<pre>1 (bases 1 to 1891) Shaul, Y.</pre>	Unpublished (1990)	luit aucomacic 2 (sites)
LOCUS DEFINITION	ACCESSION KEYWORDS SOURCE	OKGANISM	REFERENCE AUTHORS	JOURNAL	SIANDARD REFERENCE

FIG. 28 (4)

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dated 18-MAR-1991.
              Structure and function of human jun-D
                                                                                                                             175..1218
/product="junD protein"
                                                                                                               evidence=EXPERIMENTAL
                                          full staff_review
From EMBL 26 entry HSJUNDR;
                                                               Location/Qualifiers
1..1891
                                                                                                                                                                      /codon_start=1
1891..1891
Berger, I. and Shaul, Y.
                                                                                                                                                       'gene="junD"
                                                                                                 'gene="junD"
                            Unpublished (1990)
                                                                                                                                                                                    polyA_site
                                          STANDARD
                                                                                     mRNA
                            JOURNAL
AUTHORS
                                                                                                                             CDS
                                                                      FEATURES
              TITLE
                                                         COMMENT
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FIG. 28 (5)

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	CGCCAGTGGC	GGTGGCGGCA	CGCCCCCAGC	GCCGACGAGC	CGAGGGCTTC	GGCCGCTGCC	2552222225	GAGCAGCTAC	CGAACCTGTG	TGCGCTCAAG	GTTGTCGCCC	CAACCGCATC	AGAGAAAGTG	GCGCGAGCAG	GCTGCTGCCC		
	TGGGCGGCGG	TGAGTGAGCA	CCGCCGACGG	TCACCACCAC	AGGAGTTCGC	5005550505	CGGGCTCCGC	ACGCGAACCT	CCTTCGCTGC	CGCGCCTGGC	AGAGCCCGCC	AGCGGCTGCG	CGCGCCTGGA	CGAGCCTGCT	GCGGCTGCCA		
117 T	CTGAGCGGCC	ACGCTGAGCC	TACCCCCCTG	AACGGGCTGG	AGCGAGGAGC	AACCAGCTCG	GGCACGGCCA	GCGCCTGTCT	GCGACGGTCG	TTGGGGCCGC	AGCTTCGGCG	GCGGAGCGCA	GAGCGCATCT	GCGTCCACGG	CACGTCAACA		
360 G	CCATGAGGCG	GGACGCGCTG	GCCCGCCTCC	CATCCAGTCC	GGTGGCGGCC	ACACAAGCAG	GGGCCCTCG	CGCGCCCGAA	വാവാട്ടാട്ടാ	CCCAGGCGCG	CGACGTGCCG	GCGCATCAAG	GCGCAAGCTG	CACGGAGCTG	AGTCCTCAGC	CTGA	
A 405 C	CCTTCTACGG	TGATGAAGAA	CTGCGCCCGC	AGCGCCTCAT	TCTACCCCAA	TGGAGGATTT	5500500500	2552552552	252555552	CGCCGCCACC	AGACGGTGCC	ACACGCAGGA	AGTGCCGCAA	AGAGTCAGAA	TCAAGCAGAA	TCCCGGCGTA	
162	ATGGAAACAC AGCGGCGGCA	GCCGGCAGCA	GCGCTCAAGC	CCCGAGCTCG	TCACAGTTCC	GTCAAGGCCC	ອວວອວວອວວອ	GAGCTGGCCC	ອວອອວອອອວອ	CCCTTCCCGC	GACGAGCCAC	ATCGACATGG	GCCGCCTCCA	AAGACCCTCA	GTGGCGCAGC	CAGCACCAGG	
BASE COUNT ORIGIN	1	2	181 241	301	361	421	481	541	601	199	721	781	841	901	961	1021	//

FIG. 28 (6)

		154/156		
19-DEC-1991	GGCGGGATAC CACCTTGGCG TCCAGGCCCG	CGTGATCACG CGGTGGAGGT CAAAGCCCTG	CGTCTACACC CGCCGTCGGG ACCCTTTGCG AGAGGAACCG	GGCGGCCACC GAAGACACTC AGTGGCGCAG AGGGGTCAAG
19-D	CTTACGCAGC TCCTGAAACC GGGCGCGGGG	ACAGCAACGG GGGGTGGCAG ACGGTTTTGT	AGCCGCCTCC GCTCCGGGAC CACATGCACC CCGCCTTTAA	GGAACAGGCT AGGACAAGGT TAAGGGAGCA AGTTGCTGCT
159 T	CACGACGACT GACTACAAAC AAGGGTCCTG GGTCAGGGAT	ATCGTCCCCA TACCCCCGTG GGCTTTGCGG	GCTGGTCCGG CCCTCTGGAG AGCTACCTCC CGTGGCGCTT GCCACGCCGC	AAGCGGCTGC GCGCGCCTGG GCCGGTCTCC AACGGCTGCC
DNA 333 G	GCCTTTCTAT GTCTCTACAC TCGGGGTCTC CTACTTTTCG	GGAGCGCTTG ACAGTACTTT GGAGCAGGAG CCACGTGACG	GGGCGTCTAT AGCCTCTGCA GGCCACCATC GGGTTTGAGT CAGCCGCGAC	AGTGGAGCGA GGAGCGCATC GTCGAGTGCT CCATGTCAGC
1035 bp A 333 C	AAATGGAACA CTGGCAGCCT CGGATCCCTA GGGCAGGCAG	CCACGGAACT CGCCTCCGGG GCGTCACCGA ACAAGATGAA	CCGGCCCAGG GTTACTCTCC CATACCCGAC CGGCACAGGT	AGCGCATCAA AGCGGAAGCT ACGCGGGGCT AGGTCATGAC TCTGA
MUSBJUNX F 210	ATGTGCACGA GGTCGGAGCC CTCAACCTGG GAGGGCAGTG	AAGCTAGCCT ACGACGCCCA ACAGGGGGCG GACGACCTGC	GGTCCCCAGG AACCTCAGCA ACTGGGAGCT GGCGGCCACC	GAAGACCAGG AAGTGCCGGA AAGGCTGAGA CTCAAGCAGA GGACACGCCT
LOCUS BASE COUNT ORIGIN		241 301 361 421	84004	781 841 901 961 1021

FIG. 28 (7)

19-DEC-1991	TCAACGCCTC GTTCCTCCAG TAAAACAGAG CATGACCTTG GCGCCAAGAA CTCGGACCTTT CGGAGCTGGA GCGCCTGATC CCACCCAGTT CTTGTGCCCC TCGTGCGCGC CCTGGCTGAA CACAGCCGGT CAGCGGGCG GCGCGGTGG TGCCTGCAA GCAACTTCAA CCCGGGTGCG GCTGCCCTT TCCCTCGCAG GCTGCCCAACA GATCCCGGTG CCGTGCCGGA GATCCCGGGA AAAGGAAGCT GGAGCGGATC AAAAGGAAGCT GGAGCGGATC AAAAGGAAGCT GGAGCGGATC AAAAGGAAGCT GGAGCGGATC	AAGTCATGAA CCACGTTAAC TTTGA
148 T	GACGATGCCC CCTAAGATCC CCGCACCTCC ACACCGACCC GCCGAGGGCT ACCTCCGCGG GCCACCTCA GCCCACCTCA GCCCCCCCC CCCACCTCA GCCCCCCCC CCCACCTCA GCCCCCCCCCC	CTTAAGCAGA TTGCAAACGT
DNA 300 G	GACCTTCTAC CTACAGTAAC CAGTCTCAAG GCTGCTCACG TCCCAGTGTC GGCCTCAGTA TCCGGTCTAC GCCTCAGTA TCCGGTCTAC GCCTCAGTA TCCGGTCTAC GCCTCAGCG GCTCAGCCG TCCGCCTCAT	GGTGGCACAG
1005 bp A 334 C	AGATGGAAAC GTGCCTACGG ACCCGGTGGG CCGACGTCGG GCAATGGGCA ACACGCGCT CTCCCGCGGT ACAGTGAGCC GCGTGCAGCC GCGTGCAAGC GCCTGTCCCC GGCTGCAAGC	TCAGGGAACA AACTCATGCT
MUSCJUNX 223	ATGACTGCAA TCCGAGAGCG AACCTGGCCG CTCACGTCCA AAGAACGTGA CTGCATAGCC GGCATGGTGG CCGCAGCCTGC GCCAGCCTGC GCCAGCCTGC GCCAGCCTGC GCCAGCCTGC GCCAGCCTGC GCCAGCCTGC GCCAGCCTGC CTGAGCAGC CTGAGCAGCC CTGCCAGCCGC CTGAGCAGCC CTGAGCAGC	GCCAACATGC AGTGGGTGCC
LOCUS BASE COUNT ORIGIN	1 1 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1 1 2 1	901

FIG. 28 (8)

19-DEC-1991	TGGCTGCGGG TGCGTCGAGC CGCCCCGGG CCGCGCTTTC CGCTGACGCT CAGCCTGGCG CACCTTCTGC GCTGCGCCCC GGCTCCAA ACTCGCGTCG TGACCACTAC CCCGACCAGT AGGAGTTCGC CGCACGCGC CCACCCCCGG GGCCACGGGG CCGGCCCCC TGGGGCCCCC CCACCCCCGG GGCCACGCGC CCACCCCCGG GGCCACGCGC CCACCCCCGG GGCCACGCGC CCACCCCCGG GGCCACGCGC CCACCCCCGG GCCCACGCCC TCAAGACCCTC CAAATGCCGC TCAAGACCCTC CAAATGCCGC TCAAGACCCTC CAAATGCCGC AGGTGGCGCA GCTCCAAACACAG AGGTGGCGCA GCTCCAAACACAG	
129 T	CTGAGCGGCC GGTGGCTTCG AAGAAAGACG TCGGCCACTG AACGGGCTGG AGCGAGGCTGG GCCGCGCCG GCCGCTCCC CCCTTCCCGC CCCTTCCCGC CCCTTCCCGC CCCTTCCCGC CCCTTCCCGC CCCTTCCCCCC CCCTTCCCCCC CCCTTCCCCCCCC	
DNA 343 G	CGAGGAGGCG CCCCGGCGGT CAGCATGCTG GAAACCAGGG GATCCAGTCC GGTGGCAGTCC GGTGGCAGCC GGTGGCAGCC GGTGGCAGCC GCCCCCCC GAGCCAGTG ACCGCGCCTG CAAGAGGCTG CAAGAGGCTG CAAGAGGCTG	
1026 bp A. 382 C	CCTTCTATGG CTACTGGGGC CCCCGACGAGG CGGCGGGTT AGAGGCTGAT TCTACCCGAA TGGAGGCTCC ACGCCCTCC ACGCCCACCT CTCCGCATCC CTCCGCATCC ACGCCAACCT CTCCGCATCC ACGCCGACCT TGGAGCGTAT TGGAGCGTAT TGGAGCGTAT	
MUSDJUNX 172	ATGGAAACGC GTCGCTGGTG CCCGGGGCGC GACGCGCCC CCGGAGCTGG ACGCAGTTCC ACCCGGGCTC ACCCCGGTCT GCCACCGTC ACCCGGTCT ACCCCGGTCT ACCCCGGCTC ACCCGGCCAC AAAGTCCTCA AAAGTCCTCA	
LOCUS BASE COUNT ORIGIN	101 121 181 301 361 481 721 721 961	//

INTERNATIONAL SEARCH REPORT

International application No. PCT/US93/10507

A. CLASSIFICATION OF SUBJECT MATTER IPC(5) :G06F 15/42							
US CL :364/413.01 According to International Patent Classification (IPC) or to both national classification and IPC							
B. FIELDS SEARCHED							
Minimum d	ocumentation searched (classification system follow	ed by classification symbols)					
U.S. : 364/413.01; 435/6; 536/23.1							
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched							
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) MEDLINE, CA							
C. DOC	UMENTS CONSIDERED TO BE RELEVANT						
Category*	Citation of document, with indication, where a	appropriate, of the relevant passages	Relevant to claim No.				
P,X	GENETIC ENGINEERING NEWS, October 1993, Potera, "Hitachi Ch Software and Service," pages 1, 22, s	1-102					
Y	NUCLEIC ACIDS RESEARCH, Vol. 1990, Lowe et al., "A Computer Oligonucleotide Primers for Polymer 1757-1761, see entire document.	1-102					
Y	METHODS IN ENZYMOLOGY, Voi al., "Fast Alignment of DNA and Pr 502, see entire document.	1-102					
Furthe	er documents are listed in the continuation of Box (C. See patent family annex.					
	cial categories of cited documents:	*T* inter document published after the inte date and not in conflict with the applica	tion but cited to understand the				
to be part of particular relevance principle or theory underlying the invention							
	er document published on or after the international filing date ment which may throw doubts on priority claim(s) or which is	"X" document of particular relevance; the considered novel or cannot be consider when the document is taken alone	red to involve an inventive step				
cited	i to establish the publication date of another citation or other	"Y" document of particular relevance; the	claimed invention cannot be				
"O" docu	considered to involve an inventive step when the document is						
Date of the actual completion of the international search Date of mailing of the international search report							
21 Decemb	er 1993	0 3 FEB 1994					
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks		Authorized officer					
Box PCT Washington, D.C. 20231		SCOTT HOUTTEMAN A. Myza fr					
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